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# CORYNEBACTERIUM GLUTAMICUM GENES ENCODING PROTEINS INVOLVED IN HOMEOSTASIS AND ADAPTATION

## Abstract of the Disclosure

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Isolated nucleic acid molecules, designated HA nucleic acid molecules, which encode novel HA proteins from Corynebucterium glutamicum are described. The invention also provides antisense nucleic acid molecules, recombinant expression vectors containing HA nucleic acid molecules, and host cells into which the expression vectors have been introduced. The invention still further provides isolated HA proteins, mutated HA proteins, fusion proteins, antigenic peptides and methods for the improvement of production of a desired compound from C. glutamicum based on genetic engineering of HA genes in this organism.



BASF Aktiengesellschaft.

## CORYNEBACTERIUM GLUTAMICUM GENES ENCODING PROTEINS INVOLVED IN HOMEOSTASIS AND ADAPTATION

Background of the Invention

Certain products and by-products of naturally-occurring metabolic processes in cells have utility in a wide array of industries, including the food, feed, cosmetics, and pharmaceutical industries. These molecules, collectively termed 'fine chemicals', include organic acids, both proteinogenic and non-proteinogenic amino acids, nucleotides and nucleosides, lipids and fatty acids, diols, carbohydrates, aromatic 10 - compounds, vitamins and cofactors, and enzymes. Their production is most conveniently performed through the large-scale culture of bacteria developed to produce and secrete large quantities of one or more desired molecules. One particularly useful organism for this purpose is Corynebucterium glutumicum, a gram positive, nonpathogenic bacterium. Through strain selection, a number of mutant strains have been developed which produce an array of desirable compounds However, selection of strains improved for the production of a particular molecule is a time-consuming and difficult process.

Summary of the Invention

This invention provides novel nucleic acid molecules which may be used to identify or classify Corynebacterium glutamicum or related species of bacteria. C. glutamicum is a gram positive, aerobic bacterium which is commonly used in industry for the large-scale production of a variety of fine chemicals, and also for the degradation of hydrocarbons (such as in petroleum spills) and for the oxidation of terpenoids. The nucleic acid molecules therefore can be used to identify microorganisms which can be used to produce fine chemicals, e.g., by fermentation processes. While C. glutamicum itself is nonpathogenic, it is related to other Corynebacterium species, such as Corynebacterium diphtheriae (the causative agent of diphtheria), which are important human pathogens. The ability to identify the presence of Corynebacterium species therefore also can have significant clinical relevance, e.g., diagnostic applications. Further, these nucleic acid molecules may serve as reference points for the mapping of the C. glutamicum genome, or of genomes of related organisms.

These novel nucleic acid molecules encode proteins, referred to herein as homeostasis and adaptation (HA) proteins. These HA proteins are capable of, for example, performing a function involved in the maintenance of homeostasis in C. glutamicum, or in the ability of this microorganism to adapt to different environmental conditions. Given the availability of cloning vectors for use in Corynebacterium



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glutamicum, such as those disclosed in Sinskey et al., U.S. Patent No. 4,649,119, and techniques for genetic manipulation of C. glutamicum and the related Brevibacterium species (e.g., lactofermentum) (Yoshihama et al., J. Bacteriol 162: 591-597 (1985); Katsumata et al., J. Bacteriol. 159: 306-311 (1984); and Santamaria et al., J. Gen. Microbiol. 130: 2237-2246 (1984)), the nucleic acid molecules of the invention may be utilized in the genetic engineering of this organism to make it a better or more efficient producer of one or more fine chemicals. This improved production or efficiency of production of a fine chemical may be due to a direct effect of manipulation of a gene of the invention, or it may be due to an indirect effect of such manipulation.

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There are a number of mechanisms by which the alteration of an HA protein of the invention may directly affect the yield, production, and/or efficiency of production of a fine chemical from a C. glutamicum strain incorporating such an altered protein. For example, by engineering enzymes which modify or degrade aromatic or aliphatic compounds such that these enzymes are increased or decreased in activity or number, it may be possible to modulate the production of one or more fine chemicals which are the modification or degradation products of these compounds. Similarly, enzymes involved in the metabolism of inorganic compounds provide key molecules (e.g. phosphorous, sulfur, and nitrogen molecules) for the biosynthesis of such fine chemicals as amino acids, vitamins, and nucleic acids. By altering the activity or number of these enzymes in C. glutamicum, it may be possible to increase the conversion of these morganic compounds (or to use alternate inorganic compounds) to thus permit improved rates of incorporation of inorganic atoms into these fine chemicals. Genetic engineering of C. glutamicum enzymes involved in general cellular processes may also directly improve fine chemical production, since many of these enzymes directly modify fine chemicals (e.g., amino acids) or the enzymes which are involved in fine chemical synthesis or secretion. Modulation of the activity or number of cellular proteases may also have a direct effect on fine chemical production, since many proteases may degrade fine chemicals or enzymes involved in fine chemical production or breakdown.

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Further, the aforementioned enzymes which participate in aromatic/aliphatic compound modification or degradation, general biocatalysis, inorganic compound metabolism or proteolysis are each themselves fine chemicals, desirable for their activity in various in vitro industrial applications. By altering the number of copies of the gene for one or more of these enzymes in C. glutamicum it may be possible to increase the number of these proteins produced by the cell, thereby increasing the potential yield or efficiency of production of these proteins from large-scale C. glutamicum or related bacterial cultures.

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The alteration of an HA protein of the invention may also indirectly affect the yield, production, and/or efficiency of production of a fine chemical from a C glutamicum strain incorporating such an altered protein. For example, by modulating the activity and/or number of those proteins involved in the construction or rearrangement of the cell wall, it may be possible to modify the structure of the cell wall itself such that the cell is able to better withstand the mechanical and other stresses present during large-scale fermentative culture. Also, large-scale growth of C glutamicum requires significant cell wall production. Modulation of the activity or number of cell wall biosynthetic or degradative enzymes may allow more rapid rates of cell wall biosynthesis, which in turn may permit increased growth rates of this microorganism in culture and thereby increase the number of cells producing the desired fine chemical.

By modifying the HA enzymes of the invention, one may also indirectly impact the yield, production, or efficiency of production of one or more fine chemicals from C. glutamicum. For example, many of the general enzymes in C glutamicum may have a significant impact on global cellular processes (e.g., regulatory processes) which in turn have a significant effect on fine chemical metabolism. Similarly, proteases, enzymes which modify or degrade possibly toxic aromatic or aliphatic compounds, and enzymes which promote the metabolism of inorganic compounds all serve to increase the viability of C. glutamicum. The proteases aid in the selective removal of misfolded or misregulated proteins, such as those that might occur under the relatively stressful environmental conditions encountered during large-scale fermentor culture. By altering these proteins, it may be possible to further enhance this activity and to improve the viability of C glutamicum in culture. The aromatic/aliphatic modification or degradation proteins not only serve to detoxify these waste compounds (which may be encountered as impurities in culture medium or as waste products from cells themselves), but also to permit the cells to utilize alternate carbon sources if the optimal carbon source is limiting in the culture. By increasing their number and/or activity, the survival of C. glutamicum cells in culture may be enhanced. The inorganic metabolism proteins of the invention supply the cell with inorganic molecules required for all protein and nucleotide (among others) synthesis, and thus are critical for the overall viability of the cell. An increase in the number of viable cells producing one or more desired fine chemicals in large-scale culture should result in a concomitant increase in the yield, production, and/or efficiency of production of the fine chemical in the culture.

The invention provides novel nucleic acid molecules which encode proteins, referred to herein as HA proteins, which are capable of, for example, performing a function involved in the maintenance of homeostasis in C. glutamicum, or of



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participating in the ability of this microorganism to adapt to different environmental conditions. Nucleic acid molecules encoding an HA protein are referred to herein as HA nucleic acid molecules. In a preferred embodiment, an HA protein participates in C glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic

possesses a C. glutamicum enzymatic or proteolytic activity. Examples of such proteins include those encoded by the genes set forth in Table 1.

compounds, modification or degradation of aromatic or aliphatic compounds, or

Accordingly, one aspect of the invention pertains to isolated nucleic acid molecules (e.g., cDNAs) comprising a nucleotide sequence encoding an HA protein or biologically active portions thereof, as well as nucleic acid fragments suitable as primers or hybridization probes for the detection or amplification of HA-encoding nucleic acids (e.g., DNA or mRNA). In particularly preferred embodiments, the isolated nucleic acid molecule comprises one of the nucleotide sequences set forth in Appendix A or the coding region or a complement thereof of one of these nucleotide sequences. In other particularly preferred embodiments, the isolated nucleic acid molecule of the invention comprises a nucleotide sequence which hybridizes to or is at least about 50%, preferably at least about 60%, more preferably at least about 70%, 80% or 90%, and even more preferably at least about 95%, 96%, 97%, 98%, 99% or more homologous to a nucleotide sequence set forth in Appendix A, or a portion thereof. In other preferred embodiments, the isolated nucleic acid molecule encodes one of the amino acid sequences set forth in Appendix B. The preferred HA proteins of the present invention also preferably possess at least one of the HA activities described herein.

In another embodiment, the isolated nucleic acid molecule encodes a protein or portion thereof wherein the protein or portion thereof includes an amino acid sequence which is sufficiently homologous to an amino acid sequence of Appendix B, e.g., sufficiently homologous to an amino acid sequence of Appendix B such that the protein or portion thereof maintains an HA activity. Preferably, the protein or portion thereof encoded by the nucleic acid molecule maintains the ability to participate in the maintenance of homeostasis in C. glutamicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions. In one embodiment, the protein encoded by the nucleic acid molecule is at least about 50%, preferably at least about 60%, and more preferably at least about 70%, 80%, or 90% and most preferably at least about 95%, 96%, 97%, 98%, or 99% or more homologous to an amino acid sequence of Appendix B (e.g., an entire amino acid sequence selected from those sequences set forth in Appendix B). In another preferred embodiment, the protein is a full length C. glutamicum protein which is substantially homologous to an entire



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amino acid sequence of Appendix B (encoded by an open reading frame shown in Appendix A).

In another preferred embodiment, the isolated nucleic acid molecule is derived from C glutamicum and encodes a protein (e.g., an HA fusion protein) which includes a biologically active domain which is at least about 50% or more homologous to one of the amino acid sequences of Appendix B and is able to participate in the repair or recombination of DNA, in the transposition of genetic material, in gene expression (i.e., the processes of transcription or translation), in protein folding, or in protein secretion in Corynebacterium glutamicum, or has one or more of the activities set forth in Table 1, and which also includes heterologous nucleic acid sequences encoding a heterologous polypeptide or regulatory regions.

In another embodiment, the isolated nucleic acid molecule is at least 15 nucleotides in length and hybridizes under stringent conditions to a nucleic acid molecule comprising a nucleotide sequence of Appendix A. Preferably, the isolated nucleic acid molecule corresponds to a naturally-occurring nucleic acid molecule. More preferably, the isolated nucleic acid encodes a naturally-occurring *C. glutamicum* HA protein, or a biologically active portion thereof.

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Another aspect of the invention pertains to vectors, e.g., recombinant expression vectors, containing the nucleic acid molecules of the invention, and host cells into which such vectors have been introduced. In one embodiment, such a host cell is used to produce an HA protein by culturing the host cell in a suitable medium. The HA protein can be then isolated from the medium or the host cell.

Yet another aspect of the invention pertains to a genetically altered microorganism in which an HA gene has been introduced or altered. In one embodiment, the genome of the microorganism has been altered by introduction of a nucleic acid molecule of the invention encoding wild-type or mutated HA sequence as a transgene. In another embodiment, an endogenous HA gene within the genome of the microorganism has been altered, e.g., functionally disrupted, by homologous recombination with an altered HA gene. In a preferred embodiment, the microorganism belongs to the genus Corynebacterium or Brevibacterium, with Corynebacterium glutamicum being particularly preferred. In a preferred embodiment, the microorganism is also utilized for the production of a desired compound, such as an amino acid, with lysine being particularly preferred.

Still another aspect of the invention pertains to an isolated HA protein or a portion, e.g., a biologically active portion, thereof. In a preferred embodiment, the isolated HA protein or portion thereof can participate in the maintenance of homeostasis in C. glutamicum, or can perform a function involved in the adaptation of this

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microorganism to different environmental conditions. In another preferred embodiment, the isolated HA protein or portion thereof is sufficiently homologous to an amino acid sequence of Appendix B such that the protein or portion thereof maintains the ability to participate in the maintenance of homeostasis in C. glutamicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions.

The invention also provides an isolated preparation of an HA protein. In preferred embodiments, the HA protein comprises an amino acid sequence of Appendix B. In another preferred embodiment, the invention pertains to an isolated full length protein which is substantially homologous to an entire amino acid sequence of Appendix B (encoded by an open reading frame set forth in Appendix A). In yet another embodiment, the protein is at least about 50%, preferably at least about 60%, and more preferably at least about 70%, 80%, or 90% and most preferably at least about 95%, 96%, 97%, 98%, or 99% or more homologous to an entire amino acid sequence of Appendix B. In other embodiments, the isolated HA protein comprises an amino acid sequence which is at least about 50% or more homologous to one of the amino acid sequences of Appendix B and is able to participate in the maintenance of homeostasis in C glutamicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions, or has one or more of the activities set forth in Table 1.

Alternatively, the isolated HA protein can comprise an amino acid sequence which is encoded by a nucleotide sequence which hybridizes, e.g., hybridizes under stringent conditions, or is at least about 50%, preferably at least about 60%, more preferably at least about 70%, 80%, or 90%, and even more preferably at least about 95%, 96%, 97%, 98,%, or 99% or more homologous, to a nucleotide sequence of Appendix B. It is also preferred that the preferred forms of HA proteins also have one or more of the HA bioactivities described herein.

The HA polypeptide, or a biologically active portion thereof, can be operatively linked to a non-HA polypeptide to form a fusion protein. In preferred embodiments, this fusion protein has an activity which differs from that of the HA protein alone. In other preferred embodiments, this fusion protein participates in the maintenance of homeostasis in *C. glutamicum*, or performs a function involved in the adaptation of this microorganism to different environmental conditions. In particularly preferred embodiments, integration of this fusion protein into a host cell modulates production of a desired compound from the cell.

Another aspect of the invention pertains to a method for producing a fine chemical. This method involves the culturing of a cell containing a vector directing the expression of an HA nucleic acid molecule of the invention, such that a fine chemical is



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produced. In a preferred embodiment, this method further includes the step of obtaining a cell containing such a vector, in which a cell is transfected with a vector directing the expression of an HA nucleic acid. In another preferred embodiment, this method further includes the step of recovering the fine chemical from the culture. In a particularly preferred embodiment, the cell is from the genus *Corynebacterium* or *Brevibacterium*, or is selected from those strains set forth in Table 3.

Another aspect of the invention pertains to methods for modulating production of a molecule from a microorganism. Such methods include contacting the cell with an agent which modulates HA protein activity or HA nucleic acid expression such that a cell associated activity is altered relative to this same activity in the absence of the agent. In a preferred embodiment, the cell is modulated for one or more C glutamicum processes involved in cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or enzymatic or proteolytic activities. The agent which modulates HA protein activity can be an agent which stimulates HA protein activity or HA nucleic acid expression. Examples of agents which stimulate HA protein activity or HA nucleic acid expression include small molecules, active HA proteins, and nucleic acids encoding HA proteins that have been introduced into the cell. Examples of agents which inhibit HA activity or expression include small molecules and antisense HA nucleic acid molecules.

Another aspect of the invention pertains to methods for modulating yields of a desired compound from a cell, involving the introduction of a wild-type or mutant HA gene into a cell, either maintained on a separate plasmid or integrated into the genome of the host cell. If integrated into the genome, such integration can be random, or it can take place by homologous recombination such that the native gene is replaced by the introduced copy, causing the production of the desired compound from the cell to be modulated. In a preferred embodiment, said yields are increased. In another preferred embodiment, said chemical is a fine chemical. In a particularly preferred embodiment, said fine chemical is an amino acid. In especially preferred embodiments, said amino acid is L-lysine.

#### Detailed Description of the Invention

The present invention provides HA nucleic acid and protein molecules which are involved in C. glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or that have a C glutamicum enzymatic or proteolytic activity. The molecules of the invention may be utilized in the modulation of production of fine chemicals from microorganisms, such as C. glutamicum, either directly (e.g., where overexpression or



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optimization of activity of a protein involved in the production of a fine chemical (e.g., an enzyme) has a direct impact on the yield, production, and/or efficiency of production of a fine chemical from the modified C. glutamicum), or an indirect impact which nonetheless results in an increase of yield, production, and/or efficiency of production of the desired compound (e.g., where modulation of the activity or number of copies of a C. glutamicum aromatic or aliphatic modification or degradation protein results in an increase in the viability of C glutamicum cells, which in turn permits increased production in a large-scale culture setting). Aspects of the invention are further explicated below.

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#### Fine Chemicals

The term 'fine chemical' is art-recognized and includes molecules produced by an organism which have applications in various industries, such as, but not limited to, the pharmaceutical, agriculture, and cosmetics industries. Such compounds include organic acids, such as tartaric acid, itaconic acid, and diaminopimelic acid, both proteinogenic and non-proteinogenic amino acids, purine and pyrimidine bases, nucleosides, and nucleotides (as described e.g. in Kuninaka, A. (1996) Nucleotides and related compounds, p. 561-612, in Biotechnology vol. 6, Rehm et al., eds. VCH: Weinheim, and references contained therein), lipids, both saturated and unsaturated farty acids (e.g., arachidonic acid), diols (e.g., propane diol, and butane diol), carbohydrates (e.g., hyaluronic acid and trehalose), aromatic compounds (e.g., aromatic amines, vanillin, and indigo), vitamins and cofactors (as described in Ullmann's Encyclopedia of Industrial Chemistry, vol. A27, "Vitamins", p. 443-613 (1996) VCH: Weinheim and references therein; and Ong. A.S., Niki, E. & Packer, L. (1995) "Nutrition, Lipids, Health, and Disease" Proceedings of the UNESCO/Confederation of Scientific and Technological Associations in Malaysia, and the Society for Free Radical Research -Asia, held Sept. 1-3, 1994 at Penang, Malaysia, AOCS Press, (1995)), enzymes, and all other chemicals described in Gutcho (1983) Chemicals by Fermentation, Noyes Data Corporation, ISBN: 0818805086 and references therein. The metabolism and uses of certain of these fine chemicals are further explicated below.

#### A. Amino Acid Metabolism and Uses

Amino acids comprise the basic structural units of all proteins, and as such are essential for normal cellular functioning in all organisms. The term "amino acid" is art-recognized. The proteinogenic amino acids, of which there are 20 species, serve as structural units for proteins, in which they are linked by peptide bonds, while the nonproteinogenic amino acids (hundreds of which are known) are not normally found in

proteins (see Ulmann's Encyclopedia of Industrial Chemistry, vol. A2, p. 57-97 VCH: Weinheim (1985)). Amino acids may be in the D- or L- optical configuration, though L-amino acids are generally the only type found in naturally-occurring proteins. Biosynthetic and degradative pathways of each of the 20 proteinogenic amino acids have been well characterized in both prokaryotic and eukaryotic cells (see, for example, Stryer, L. Biochemistry, 3<sup>rd</sup> edition, pages 578-590 (1988)). The 'essential' amino acids (histidine, isoleucine, leucine, lysine, methionine, phenylalanine, threonine, tryptophan, and valine), so named because they are generally a nutritional requirement due to the complexity of their biosynthesis, are readily converted by simple biosynthetic pathways to the remaining 11 'nonessential' amino acids (alanine, arginine, asparagine, aspartate, cysteine, glutamate, glutamine, glycine, proline, serine, and tyrosine). Higher animals do retain the ability to synthesize some of these amino acids, but the essential amino acids must be supplied from the diet in order for normal protein synthesis to occur.

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Aside from their function in protein biosynthesis, these amino acids are interesting chemicals in their own right, and many have been found to have various applications in the food, feed, chemical, cosmetics, agriculture, and pharmaceutical industries. Lysine is an important amino acid in the nutrition not only of humans, but also of monogastric animals such as poultry and swine. Glutamate is most commonly used as a flavor additive (mono-sodium glutamate, MSG) and is widely used throughout the food industry, as are aspartate, phenylalanine, glycine, and cysteine. Glycine, Lmethionine and tryptophan are all utilized in the pharmaceutical industry. Glutamine, valine, leucine, isoleucine, histidine, arginine, proline, serine and alanine are of use in both the pharmaceutical and cosmetics industries. Threonine, tryptophan, and D/ Lmethionine are common feed additives. (Leuchtenberger, W. (1996) Amino aids technical production and use, p. 466-502 in Rehm et al. (eds.) Biotechnology vol. 6, chapter 14a, VCH. Weinheim). Additionally, these amino acids have been found to be useful as precursors for the synthesis of synthetic amino acids and proteins, such as Nacetylcysteine, S-carboxymethyl-L-cysteine, (S)-5-hydroxytryptophan, and others described in Ulmann's Encyclopedia of Industrial Chemistry, vol. A2, p. 57-97, VCH: Weinheim, 1985.

The biosynthesis of these natural amino acids in organisms capable of producing them, such as bacteria, has been well characterized (for review of bacterial amino acid biosynthesis and regulation thereof, see Umbarger, H.E.(1978) Ann Rev. Biochem. 47: 533-606). Glutamate is synthesized by the reductive amination of a-ketoglutarate, an intermediate in the citric acid cycle. Glutamine, proline, and arginine are each subsequently produced from glutamate. The biosynthesis of serine is a three-step process beginning with 3-phosphoglycerate (an intermediate in glycolysis), and

resulting in this amino acid after oxidation, transamination, and hydrolysis steps. Both cysteine and glycine are produced from serine; the former by the condensation of homocysteine with serine, and the latter by the transferal of the side-chain  $\beta$ -carbon atom to tetrahydrofolate, in a reaction catalyzed by serine transhydroxymethylase.

5 Phenylalanine, and tyrosine are synthesized from the glycolytic and pentose phosphate pathway precursors erythrose 4-phosphate and phosphoenolpyruvate in a 9-step biosynthetic pathway that differ only at the final two steps after synthesis of prephenate. Tryptophan is also produced from these two initial molecules, but its synthesis is an 11step pathway. Tyrosine may also be synthesized from phenylalanine, in a reaction catalyzed by phenylalanine hydroxylase. Alanine, valine, and leucine are all biosynthetic products of pyruvate, the final product of glycolysis. Aspartate is formed from oxaloacetate, an intermediate of the citric acid cycle. Asparagine, methionine, threonine, and lysine are each produced by the conversion of aspartate. Isoleucine is formed from threonine. A complex 9-step pathway results in the production of histidine from 5-phosphoribosyl-1-pyrophosphate, an activated sugar.

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Amino acids in excess of the protein synthesis needs of the cell cannot be stored, and are instead degraded to provide intermediates for the major metabolic pathways of the cell (for review see Stryer, L. Biochemistry 3rd ed. Ch. 21 "Amino Acid Degradation and the Urea Cycle" p. 495-516 (1988)). Although the cell is able to convert unwanted amino acids into useful metabolic intermediates, amino acid production is costly in terms of energy, precursor molecules, and the enzymes necessary to synthesize them. Thus it is not surprising that amino acid biosynthesis is regulated by feedback inhibition, in which the presence of a particular amino acid serves to slow or entirely stop its own -production (for overview of feedback mechanisms in amino acid biosynthetic pathways, see Stryer, L. Biochemistry, 3rd ed. Ch. 24: "Biosynthesis of Amino Acids and Heme" p. 575-600 (1988)). Thus, the output of any particular amino acid is limited by the amount of that amino acid present in the cell.

### B. Vitamin, Cofactor, and Nutruceutical Metabolism and Uses

Vitamins, cofactors, and nutraceuticals comprise another group of molecules which the higher animals have lost the ability to synthesize and so must ingest, although they are readily synthesized by other organisms such as bacteria. These molecules are either bioactive substances themselves, or are precursors of biologically active substances which may serve as electron carriers or intermediates in a variety of metabolic pathways. Aside from their nutritive value, these compounds also have significant industrial value as coloring agents, antioxidants, and catalysts or other processing aids. (For an overview of the structure, activity, and industrial applications

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of these compounds, see, for example, Ullman's Encyclopedia of Industrial Chemistry, "Vitamins" vol. A27, p. 443-613, VCH: Weinheim, 1996.) The term "vitamin" is arrrecognized, and includes nutrients which are required by an organism for normal functioning, but which that organism cannot synthesize by itself. The group of vitamins may encompass cofactors and nutraceutical compounds. The language "cofactor" includes nonproteinaceous compounds required for a normal enzymatic activity to occur. Such compounds may be organic or inorganic; the cofactor molecules of the invention are preferably organic. The term "nutraceutical" includes dietary supplements having health benefits in plants and animals, particularly humans. Examples of such molecules are vitamins, antioxidants, and also certain lipids (e.g., polyunsaturated fatty acids).

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The biosynthesis of these molecules in organisms capable of producing them. such as bacteria, has been largely characterized (Ullman's Encyclopedia of Industrial Chemistry, "Vitamins" vol. A27, p. 443-613, VCH Weinheim, 1996; Michal, G. (1999) Biochemical Pathways: An Atlas of Biochemistry and Molecular Biology, John Wiley & Sons; Ong, A.S., Niki, E. & Packer, L. (1995) "Nutrition, Lipids, Health, and Disease" Proceedings of the UNESCO/Confederation of Scientific and Technological Associations in Malaysia, and the Society for Free Radical Research – Asia, held Sept. 1-3, 1994 at Penang, Malaysia, AOCS Press: Champaign, IL X, 374 S).

Thiamin (vitamin B<sub>1</sub>) is produced by the chemical coupling of pyrimidine and thiazole moieties. Riboflavin (vitamin B2) is synthesized from guanosine-5'-triphosphate (GTP) and ribose-5'-phosphate. Ribotlavin, in turn, is utilized for the synthesis of flavin mononucleotide (FMN) and flavin adenine dinucleotide (FAD). The family of compounds collectively termed 'vitamin B6' (e.g., pyridoxine, pyridoxamine, pyridoxa-5'-phosphate, and the commercially used pyridoxin hydrochloride) are all derivatives of the common structural unit, 5-hydroxy-6-methylpyridine. Pantothenate (pantothenic acid, (R)-(+)-N-(2,4-dihydroxy-3,3-dimethyl-1-oxobutyl)- $\beta$ -alanine) can be produced either by chemical synthesis or by fermentation. The final steps in pantothenate biosynthesis consist of the ATP-driven condensation of  $\beta$ -alanine and pantoic acid. The enzymes responsible for the biosynthesis steps for the conversion to pantoic acid, to  $\beta$ alanine and for the condensation to panthotenic acid are known. The metabolically active form of pantothenate is Coenzyme A, for which the biosynthesis proceeds in 5 enzymatic steps. Pantothenate, pyridoxal-5'-phosphate, cysteine and ATP are the precursors of Coenzyme A. These enzymes not only catalyze the formation of panthothante, but also the production of (R)-pantoic acid, (R)-pantolacton, (R)panthenol (provitamin B<sub>5</sub>), pantetheine (and its derivatives) and coenzyme A.

Biotin biosynthesis from the precursor molecule pimeloyl-CoA in microorganisms has been studied in detail and several of the genes involved have been identified. Many of the corresponding proteins have been found to also be involved in Fe-cluster synthesis and are members of the nifS class of proteins. Lipoic acid is derived from octanoic acid, and serves as a coenzyme in energy metabolism, where it becomes part of the pyruvate dehydrogenase complex and the  $\alpha$ -ketoglutarate dehydrogenase complex. The folates are a group of substances which are all derivatives of folic acid, which is turn is derived from L-glutamic acid, p-amino-benzoic acid and 6methylpterin. The biosynthesis of folic acid and its derivatives, starting from the metabolism intermediates guanosine-5'-triphosphate (GTP), L-glutamic acid and pamino-benzoic acid has been studied in detail in certain microorganisms.

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.Corrinoids (such as the cobalamines and particularly vitamin B<sub>12</sub>) and porphyrines belong to a group of chemicals characterized by a tetrapyrole ring system The biosynthesis of vitamin B<sub>12</sub> is sufficiently complex that it has not yet been 15 completely characterized, but many of the enzymes and substrates involved are now known. Nicotinic acid (nicotinate), and nicotinamide are pyridine derivatives which are also termed 'niacin' Niacin is the precursor of the important coenzymes NAD (nicotinamide adenine dinucleotide) and NADP (nicotinamide adenine dinucleotide phosphate) and their reduced forms.

The large-scale production of these compounds has largely relied on cell-free chemical syntheses, though some of these chemicals have also been produced by largescale culture of microorganisms, such as riboflavin, Vitamin B6, pantothenate, and biotin. Only Vitamin B<sub>12</sub> is produced solely by fermentation, due to the complexity of its synthesis. In vitro methodologies require significant inputs of materials and time, often at great cost.

## C. Purine, Pyrimidine, Nucleoside and Nucleotide Metabolism and Uses

Purine and pyrimidine metabolism genes and their corresponding proteins are important targets for the therapy of tumor diseases and viral infections. The language "purine" or "pyrimidine" includes the nitrogenous bases which are constituents of nucleic acids, co-enzymes, and nucleotides. The term "nucleotide" includes the basic structural units of nucleic acid molecules, which are comprised of a nitrogenous base, a pentose sugar (in the case of RNA, the sugar is ribose; in the case of DNA, the sugar is D-deoxyribose), and phosphoric acid. The language "nucleoside" includes molecules which serve as precursors to nucleotides, but which are lacking the phosphoric acid **35**、 moiety that nucleotides possess. By inhibiting the biosynthesis of these molecules, or their mobilization to form nucleic acid molecules, it is possible to inhibit RNA and DNA

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synthesis; by inhibiting this activity in a fashion targeted to cancerous cells, the ability of tumor cells to divide and replicate may be inhibited. Additionally, there are nucleotides which do not form nucleic acid molecules, but rather serve as energy stores (i.e., AMP) or as coenzymes (i.e., FAD and NAD).

Several publications have described the use of these chemicals for these medical indications, by influencing purine and/or pyrimidine metabolism (e.g. Christopherson, R.I. and Lyons, S.D. (1990) "Potent inhibitors of de novo pyrimidine and purine biosynthesis as chemotherapeutic agents." Med Res Reviews 10: 505-548). Studies of enzymes involved in purine and pyrimidine metabolism have been focused on the development of new drugs which can be used, for example, as immunosuppressants or anti-proliferants (Smith, J.L., (1995) "Enzymes in nucleotide synthesis." Curr. Opin. Struct. Biol 5: 752-757; (1995) Riochem Soc Transact. 23: 877-902). However, purine and pyrimidine bases, nucleosides and nucleotides have other utilities: as intermediates in the biosynthesis of several fine chemicals (e.g., thiamine, S-adenosyl-methionine, 15 folates, or riboflavin), as energy carriers for the cell (e.g., ATP or GTP), and for chemicals themselves, commonly used as flavor enhancers (e.g., IMP or GMP) or for several medicinal applications (see, for example, Kuninaka, A. (1996) Nucleotides and Related Compounds in Biotechnology vol. 6, Rehm et al., eds. VCH: Weinheim, p. 561-612). Also, enzymes involved in purine, pyrimidine, nucleoside, or nucleotide metabolism are increasingly serving as targets against which chemicals for crop protection, including fungicides, herbicides and insecticides, are developed.

The metabolism of these compounds in bacteria has been characterized (for reviews see, for example, Zalkin, H. and Dixon, J.E. (1992) "de novo purine nucleotide" biosynthesis", in: Progress in Nucleic Acid Research and Molecular Biology, vol. 42, Academic Press:, p. 259-287; and Michal, G. (1999) "Nucleotides and Nucleosides", Chapter 8 in: Biochemical Pathways: An Atlas of Biochemistry and Molecular Biology, Wiley: New York). Purine metabolism has been the subject of intensive research, and is essential to the normal functioning of the cell. Impaired purine metabolism in higher animals can cause severe disease, such as gout. Purine nucleotides are synthesized from 30 ribose-5-phosphate, in a series of steps through the intermediate compound inosine-5'phosphate (IMP), resulting in the production of guanosine-5'-monophosphate (GMP) or adenosine-5'-monophosphate (AMP), from which the triphosphate forms utilized as nucleotides are readily formed. These compounds are also utilized as energy stores, so their degradation provides energy for many different biochemical processes in the cell. Pyrimidine biosynthesis proceeds by the formation of uridine-5'-monophosphate (UMP) from ribose-5-phosphate. UMP, in turn, is converted to cytidine-5'-triphosphate (CTP). The deoxy- forms of all of these nucleotides are produced in a one step reduction

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reaction from the diphosphate ribose form of the nucleotide to the diphosphate deoxyribose form of the nucleotide. Upon phosphorylation, these molecules are able to participate in DNA synthesis.

#### D. Trehalose Metabulism and Uses

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Trehalose consists of two glucose molecules, bound in a, a-1,1 linkage. It is commonly used in the food industry as a sweetener, an additive for dried or frozen foods, and in beverages. However, it also has applications in the pharmaceutical, cosmetics and biotechnology industries (see, for example, Nishimoto et al., (1998) U.S. Patent No. 5,759,610; Singer, M.A. and Lindquist, S. (1998) Trends Biotech. 16: 460-467; Paiva, C.L.A. and Panek, A.D. (1996) Biotech. Ann. Rev. 2. 293-314; and Shiosaka, M. (1997) J. Japan 172: 97-102). Trehalose is produced by enzymes from many microorganisms and is naturally released into the surrounding medium, from which it can be collected using methods known in the art.

Maintenance of Homeostasis in C. glutamicum and Environmental Adaptation

The metabolic and other biochemical processes by which cells function are sensitive to environmental conditions such as temperature, pressure, solute concentration, and availability of oxygen. When one or more such environmental 20 condition is perturbed or altered in a fashion that is incompatible with the normal functioning of these cellular processes, the cell must act to maintain an intracellular environment which will permit them to occur despite the hostile extracellular environment. Gram positive bacterial cells, such as C. glutamicum cells, have a number of mechanisms by which internal homeostasis may be maintained despite unfavorable extracellular conditions. These include a cell wall, proteins which are able to degrade possibly toxic aromatic and aliphatic compounds, mechanisms of proteolysis whereby misfolded or misregulated proteins may be rapidly destroyed, and catalysts which permit intracellular reactions to occur which would not normally take place under the conditions optimal for bacterial growth.

Aside from merely surviving in a hostile environment, bacterial cells (e.g. C glutamicum cells) are also frequently able to adapt such that they are able to take advantage of such conditions. For example, cells in an environment lacking desired carbon sources may be able to adapt to growth on a less-suitable carbon source Also, cells may be able to utilize less desirable inorganic compounds when the commonly utilized ones are unavailable. C. glutumicum cells possess a number of genes which permit them to adapt to utilize inorganic and organic molecules which they would normally not encounter under optimal growth conditions as nutrients and precursors for

metabolism. Aspects of cellular processes involved in homeostasis and adaptation are further explicated below.

A Modification and Degradation of Aromatic and Aliphatic Compounds

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Bacterial cells are routinely exposed to a variety of aromatic and aliphatic compounds in nature. Aromatic compounds are organic molecules having a cyclic ring structure, while aliphatic compounds are organic molecules having open chain structures rather than ring structures. Such compounds may arise as by-products of industrial processes (e.g., benzene or toluene), but may also be produced by certain microorganisms (e.g., alcohols). Many of these compounds are toxic to cells, particularly the aromatic compounds, which are highly reactive due to the high-energy ring structure. Thus, certain bacteria have developed mechanisms by which they are able to modify or degrade these compounds such that they are no longer hazardous to the cell. Cells may possess enzymes that are able to, for example, hydroxylate, isomerize, or methylate aromatic or aliphatic compounds such that they are either rendered less toxic, or such that the modified form is able to be processed by standard cellular waste and degradation pathways. Also, cells may possess enzymes which are able to specifically degrade one or more such potentially hazardous substance, thereby protecting the cell. Principles and examples of these types of modification and degradation processes in bacteria are described in several publications, e.g., Sahm, H. (1999) "Procaryotes in Industrial Production" in Lengeler, J.W. et al., eds. Biology of the Procaryotes, Thieme Verlag: Stuttgart; and Schlegel, H.G. (1992) Allgemeine Mikrobiologie, Thieme: Stuttgart).

Aside from simply inactivating hazardous aromatic or aliphatic compounds, many bacteria have evolved to be able to utilize these compounds as carbon sources for continued metabolism when the preferred carbon sources of the cell are not available. For example, Pseudomonas strains able to utilize toluene, benzene, and 1,10-dichlorodecane as carbon sources are known (Chang, B.V. et al. (1997) Chemosphere 35(12): 2807-2815; Wischnak, C. et al. (1998) Appl Environ. Microbiol. 64(9): 3507-3511; Churchill, S.A. et al. (1999) Appl Environ Microbiol. 65(2): 549-552). There are similar examples from many other bacterial species which are known in the art.

The ability of certain bacteria to modify or degrade aromatic and aliphatic compounds has begun to be exploited. Petroleum is a complex mixture of chemicals which includes aliphatic molecules and aromatic compounds. By applying bacteria having the ability to degrade or modify these toxic compounds to an oil spill, for example, it is possible to eliminate much of the environmental damage with high efficiency and low cost (see, for example, Smith, M.R. (1990) "The biodegradation of aromatic hydrocarbons by bacteria"

Biodegradation 1(2-3): 191-206; and Suyama, T. et al. (1998) "Bacterial isolates degrading aliphatic polycarbonates," FEMS Microbiol Lett. 161(2): 255-261).

## B Metabolism of Inorganic Compounds

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Cells (e.g., bacterial cells) contain large quantities of different molecules, such as water, inorganic ions, and organic substances (e.g., proteins, sugars, and other macromolecules). The bulk of the mass of a typical cell consists of only 4 types of atoms: carbon, oxygen, hydrogen, and nitrogen. Although they represent a smaller percentage of the content of a cell, inorganic substances are equally as important to the proper functioning of the cell. Such molecules include phosphorous, sulfur, calcium, magnesium, iron, zinc, manganese, copper, molybdenum, tungsten, and cobalt. Many of these compounds are critical for the construction of important molecules, such as nucleotides (phosphorous) and amino acids (nitrogen and sulfur). Others of these inorganic ions serve as cofactors for enzymic reactions or contribute to osmotic pressure. All such molecules must be taken up by the bacterium from the surrounding environment.

For each of these inorganic compounds it is desirable for the bacterium to take up the form which can be most readily used by the standard metabolic machinery of the cell. However, the bacterium may encounter environments in which these preferred forms are not readily available. In order to survive under these circumstances, it is important for bacteria to have additional biochemical mechanisms which are able to convert less metabolically active but readily available forms of these inorganic compounds to ones which may be used in cellular metabolism. Bacteria frequently possess a number of genes encoding enzymes for this purpose, which are not expressed unless the desired inorganic species are not available. Thus, these genes for the metabolism of various inorganic compounds serve as another tool which bacteria may use to adapt to suboptimal environmental conditions.

After carbon, the most important element in the cell is nitrogen. A typical bacterial cell contains between 12-15% nitrogen. It is a constituent of amino acids and nucleotides, as well as many other important molecules in the cell. Further, nitrogen may serve as a substitute for oxygen as a terminal electron acceptor in energy metabolism. Good sources of nitrogen include many organic and inorganic compounds, such ammonia gas or ammonia salts (e.g., NH<sub>4</sub>Cl, (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, or NH<sub>4</sub>OH), nitrates, urea, amino acids, or complex nitrogen sources like corn steep liquor, soy bean flour, soy bean protein, yeast extract, meat extract, etc. Ammonia nitrogen is fixed by the action of particular enzymes: glutamate dehydrogenase, glutamine synthase, and glutamine-2-oxoglutarate aminotransferase. The transfer of amino-nitrogen from one organic molecule to another is accomplished by the aminotransferases, a class of enzymes which transfer one amino group from an alpha-amino acid to an alpha-keto acid. Nitrate may be reduced via nitrate reductase, nitrite reductase,

and further redox enzymes until it is converted to molecular nitrogen or ammonia, which may be readily utilized by the cell in standard metabolic pathways.

Phosphorous is typically found intracellularly in both organic and inorganic forms, and may be taken up by the cell in either of these forms as well, though most microorganisms preferentially take up inorganic phosphate. The conversion of organic phosphate to a form which the cell can utilize requires the action of phosphatases (e.g., phytases, which hydrolyze phyate-yielding phosphate and inositol derivatives). Phosphate is a key element in the synthesis of nucleic acids, and also has a significant role in cellular energy metabolism (e.g., in the synthesis of ATP, ADP, and AMP).

Sulfur is a requirement for the synthesis of amino acids (e.g., methionine and cysteine), vitamins (e.g., thiamine, biotin, and lipoic acid) and iron sulfur proteins. Bacteria obtain sulfur primarily from inorganic sulfate, though thiosulfate, sulfite, and sulfide are also commonly utilized. Under conditions where these compounds may not be readily available, many bacteria express genes which enable them to utilize sulfonate compounds such as 2-aminosulfonate (taurine) (Kertesz, M.A. (1993) "Proteins induced by sulfate limitation in Escherichia coti, Pseudomonas putida, or Staphylococcus aureus." J Bacteriol. 175: 1187-1190).

Other inorganic atoms, e.g., metal or calcium ions, are also critical for the viability of cells. Iron, for example, plays a key role in redox reactions and is a cofactor of iron-sulfur proteins, heme proteins, and cytochromes. The uptake of iron into bacterial cells may be accomplished by the action of siderophores, chelating agents which bind extracellular iron ions and translocate them to the interior of the cell. For reference on the metabolism of iron and other inorganic compounds, see: Lengeler et al. (1999) Biology of Prokaryotes, Thieme Verlag: Stuttgart; Neidhardt, F.C. et al., eds. Escherichia coli and Salmonella. ASM Press: Washington, D.C.; Sonenshein, A.L. et al., eds. (199?) Bacillus subrilis and Other Gram-Positive Bacteria, ASM Press: Washington, D.C.; Voet, D. and Voet, J.G. (1992) Biochemie, VCH: Weinheim; Brock, T.D. and Madigan, M.T. (1991) Biology of Microorgansisms, 6th ed. Prentice Hall: Englewood Cliffs, p. 267-269; Rhodes, P.M. and Stanbury, P.F. Applied Microbial Physiology - A Practical Approach, Oxford Univ. Press: Oxford. 30

#### C. Enzymes and Proteolysis

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The intracellular conditions for which bacteria such as C glutamicum are optimized are frequently not conditions under which many biochemical reactions would normally take place. In order to make such reactions proceed under physiological conditions, cells utilize enzymes. Enzymes are proteinaceous biological catalysts, spatially orienting reacting molecules or providing a specialized environment such that

the energy barrier to a biochemical reaction is lowered. Different enzymes catalyze different reactions, and each enzyme may be the subject of transcriptional, translational, or posttranslational regulation such that the reaction will only take place under appropriate conditions and at specified times. Enzymes may contribute to the degradation (e.g., the proteases), synthesis (e.g., the synthases), or modification (e.g., transferases or isomerases) of compounds, all of which enable the production of necessary compounds within the cell. This, in turn, contributes to the maintenance of cellular homeostasis.

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However, the fact that enzymes are optimized for activity under the physiological conditions at which the bacterium is most viable means that when environmental conditions are perturbed, there is a significant possibility that enzyme activity will also be perturbed. For example, changes in temperature may result in aberrantly folded proteins, and the same is true for changes of pH - protein folding is largely dependent on electrostatic and hydrophobic interactions of amino acids within the polypeptide chain, so any alteration to the charges on individual amino acids (as might be brought about by a change in cellular pH) may have a profound effect on the ability of the protein to correctly fold. Changes in temperature effectively change the amount of kinetic energy that the polypeptide molecule possesses, which affects the ability of the polypeptide to settle into a correctly folded, energetically stable configuration. Misfolded proteins may be harmful to the cell for two reasons. First, the aberrantly folded protein may have a similarly aberrant activity, or no activity whatsoever. Second, misfolded proteins may lack the conformational regions necessary for proper regulation by other cellular systems and thus may continue to be active but in an uncontrolled fashion.

The cell has a mechanism by which mistolded enzymes and regulatory proteins may be rapidly destroyed before any damage occurs to the cell: proteolysis. Proteins such as those of the la/lon family and those of the Clp family specifically recognize and degrade misfolded proteins (see, e.g., Sherman, M.Y., Goldberg, A.L. (1999) EXS 77: 57-78 and references therein and Porankiewicz J. (1999) Molec. Microbiol. 32(3): 449-58, and references therein; Neidhardt, F.C., et al. (1996) E. coli and Salmonella, ASM Press: Washington, D.C. and references therein; and Pritchard, G.G., and Coolbear, T. (1993) FEMS Microbiol. Rev. 12(1-3): 179-206 and references therein). These enzymes bind to misfolded or unfolded proteins and degrade them in an ATP-dependent manner. Proteolysis thus serves as an important mechanism employed by the cell to prevent damage to normal cellular functions upon environmental changes, and it further permits cells to survive under conditions and in environments which would otherwise be toxic due to misregulated and/or aberrant enzyme or regulatory activity.

Proteolysis also has important functions in the cell under optimal environmental conditions. Within normal metabolic processes, proteases aid in the hydrolysis of peptide bonds, in the catabolism of complex molecules to provide necessary degradation products, and in protein modification. Secreted proteases play an important role in the catabolism of external nutrients even prior to the entry of these compounds into the cell. Further, proteolytic activity itself may serve regulatory functions; sporulation in 8 subults and cell cycle progression in Caulobucter spp. are known to be regulated by key proteolytic events in each of these species (Gottesman, S. (1999) Curr. Opin. Microbiol. 2(2): 142-147). Thus, proteolytic processes are key for cellular survival under both suboptimal and optimal environmental conditions, and contribute to the overall maintenance of homeostasis in cells.

#### D. Cell Wall Production and Rearrangements

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While the biochemical machinery of the cell may be able to readily adapt to different and possibly unfavorable environments, cells still require a general mechanism by which they may be protected from the environment. For many bacteria, the cell wall affords such protection, and also plays roles in adhesion, cell growth and division, and transport of desired solutes and waste materials.

In order to function, cells require intracellular concentrations of metabolites and other molecules that are substantially higher than those of the surrounding media. Since these metabolites are largely prevented from leaving the cell due to the presence of the hydrophobic membrane, the tendency of the system is for water molecules to enter the cell from the external medium such that the interior concentrations of solutes match the exterior concentrations. Water molecules are readily able to cross the cellular membrane, and this membrane is not able to withstand the resulting swelling and pressure, which may lead to osmotic lysis of the cell. The rigidity of the cell wall greatly improves the ability of the cell to tolerate these pressures, and offers a further barrier to the unwanted diffusion of these metabolites and desired solutes from the cell. Similarly, the cell wall also serves to prevent unwanted material from entering the cell.

The cell wall also participates in a number of other cellular processes, such as adhesion and cell growth and division. Due to the fact that the cell wall completely surrounds the cell, any interaction of the cell with its surroundings must be mediated by the cell wall. Thus, the cell wall must participate in any adherence of the cell to other cells and to desired surfaces. Further, the cell cannot grow or divide without concomitant changes in the cell wall. Since the protection that the wall affords requires its presence during growth, morphogenesis and multiplication, one of the key steps in cell division is cell wall synthesis within the cell such that a new cell divides from the old. Thus, frequently cell wall

biosynthesis is regulated in tandem with cell growth and cell division (see, e.g., Sonenshein, A.L. et al, eds. (1993) *Bucillus subtilis* and Other Gram-Positive Bacteria, ASM: Washington, D.C.).

The structure of the cell wall varies between gram-positive and gram-negative bacteria. However, in both types, the fundamental structural unit of the wall remains similar: an overlapping lattice of two polysaccharides, N-acetyl glucosamine (NAG) and N-acetyl muramic acid (NAM) which are cross-linked by amino acids (most commonly L-alanine, D-glutamate, diaminopimelic acid, and D-alanine), termed 'peptidoglycan'. The processes involved in the synthesis of the cell wall are known (see, e.g., Michal, G., ed. (1999) Biochemical Pathways: An Atlas of Biochemistry and Molecular Biology, Wiley: New York).

In gram-negative baoteria, the inner cellular membrane is coated by a single-layered peptidoglycan (approximately 10 nm thick), termed the murein-sacculus. This peptidoglycan structure is very rigid, and its structure determines the shape of the organism. The outer surface of the murein-sacculus is covered with an outer membrane, containing porins and other membrane proteins, phospholipids, and lipopolysaccharides. To maintain a tight association with the outer membrane, the gram-negative cell wall also has interspersed lipid molecules which serve to anchor it to the surrounding membrane.

In gram-positive bacteria, such as Corynebucterium glutamicum, the cytoplasmic membrane is covered by a multi-layered peptidoglycan, which ranges from 20-80 nm in thickness (see, e.g., Lengeler et al. (1999) Biology of Prokaryotes Thieme Verlag: Stuttgart, p. 913-918, p. 875-899, and p. 88-109 and references therein). The gram-positive cell wall also contains teichoic acid, a polymer of glycerol or ribitol linked through phosphate groups. Teichoic acid is also able to associate with amino acids, and forms covalent bonds with muramic acid. Also present in the cell wall may be lipoteichoic acids and teichuronic acids. If present, cellular surface structures such as flagella or capsules will be anchored in this layer as well.

#### III. Elements and Methods of the Invention

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The present invention is based, at least in part, on the discovery of novel molecules, referred to herein as HA nucleic acid and protein molecules, which participate in the maintenance of homeostasis in C glutamicum, or which perform a function involved in the adaptation of this microorganism to different environmental conditions. In one embodiment, the HA molecules participate in C glutamicum cell wall biosynthesis or rearrangements, in the metabolism of inorganic compounds, in the modification or degradation of aromatic or aliphatic compounds, or have an enzymatic or proteolytic activity. In a preferred embodiment, the activity of the HA molecules of

the present invention with regard to C glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or enzymatic or proteolytic activity has an impact on the production of a desired fine chemical by this organism. In a particularly preferred embodiment, the HA molecules of the invention are modulated in activity, such that the C glutamicum cellular processes in which the HA molecules participate (e.g., C glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or enzymatic or proteolytic activity) are also altered in activity, resulting either directly or indirectly in a modulation of the yield, production, and/or efficiency of production of a desired fine chemical by C glutamicum.

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The language, "HA protein" or "HA polypeptide" includes proteins which participate in a number of cellular processes related to C. gluiamicum homeostasis or the ability of C glutamicum cells to adapt to unfavorable environmental conditions. For example, an HA protein may be involved in C. glutamicum cell wall biosynthesis or rearrangements, in the metabolism of inorganic compounds in C glutamicum, in the modification or degradation of aromatic or aliphatic compounds in C. glutamicum, or have a C. glutamicum enzymatic or proteolytic activity. Examples of HA proteins include those encoded by the HA genes set forth in Table 1 and Appendix A. The terms "HA gene" or "HA nucleic acid sequence" include nucleic acid sequences encoding an HA protein, which consist of a coding region and also corresponding untranslated 5' and 3' sequence regions. Examples of HA genes include those set forth in Table 1. The terms "production" or "productivity" are art-recognized and include the concentration of the fermentation product (for example, the desired fine chemical) formed within a given time and a given fermentation volume (e.g., kg product per hour per liter). The term "efficiency of production" includes the time required for a particular level of production to be achieved (for example, how long it takes for the cell to attain a particular rate of output of a fine chemical). The term "yield" or "product/carbon yield" is art-recognized and includes the efficiency of the conversion of the carbon source into the product (i.e., fine chemical). This is generally written as, for example, kg product per kg carbon source. By increasing the yield or production of the compound, the quantity of recovered molecules, or of useful recovered molecules of that compound in a given amount of culture over a given amount of time is increased. The terms "biosynthesis" or a "biosynthetic pathway" are art-recognized and include the synthesis of a compound, preferably an organic compound, by a cell from intermediate compounds in what may be a multistep and highly regulated process. The terms "degradation" or a "degradation pathway" are art-recognized and include the breakdown of a compound, preferably an

organic compound, by a cell to degradation products (generally speaking, smaller or less complex molecules) in what may be a multistep and highly regulated process. The language "metabolism" is art-recognized and includes the totality of the biochemical reactions that take place in an organism. The metabolism of a particular compound, then, (e.g., the metabolism of an amino acid such as glycine) comprises the overall biosynthetic, modification, and degradation pathways in the cell related to this compound. The term "homeostasis" is art-recognized and includes all of the mechanisms utilized by a cell to maintain a constant intracellular environment despite the prevailing extracellular environmental conditions. A non-limiting example of such processes is the utilization of a cell wall to prevent osmotic lysis due to high intracellular solute concentrations. The term "adaptation" or "adaptation to an environmental condition" is an-recognized and includes mechanisms utilized by the cell to render the cell able to survive under nonpreferred environmental conditions (generally speaking, those environmental conditions in which one or more favored nutrients are absent, or in which an environmental condition such as temperature, pH, osmolarity, oxygen percentage and the like fall outside of the optimal survival range of the cell). Many cells, including C glutamicum cells, possess genes encoding proteins which are expressed under such environmental conditions and which permit continued growth in such suboptimal conditions.

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In another embodiment, the HA molecules of the invention are capable of **20** modulating the production of a desired molecule, such as a fine chemical, in a microorganism such as C. glutamicum. There are a number of mechanisms by which the alteration of an HA protein of the invention may directly affect the yield, production, and/or efficiency of production of a fine chemical from a C. glutamicum strain incorporating such an altered protein. For example, by engineering enzymes which 25 modify or degrade aromatic or aliphatic compounds such that these enzymes are increased or decreased in activity or number, it may be possible to modulate the production of one or more fine chemicals which are the modification or degradation products of these compounds Similarly, enzymes involved in the metabolism of inorganic compounds provide key molecules (e.g. phosphorous, sulfur, and nitrogen molecules) for the biosynthesis of such fine chemicals as amino acids, vitamins, and nucleic acids. By altering the activity or number of these enzymes in C. glutamicum, it may be possible to increase the conversion of these inorganic compounds (or to use alternate inorganic compounds) to thus permit improved rates of incorporation of inorganic atoms into these fine chemicals. Genetic engineering of C. glutamicum 35 enzymes involved in general cellular processes may also directly improve fine chemical production, since many of these enzymes directly modify fine chemicals (e.g., amino

acids) or the enzymes which are involved in fine chemical synthesis or secretion. Modulation of the activity or number of cellular proteases may also have a direct effect on fine chemical production, since many proteases may degrade fine chemicals or enzymes involved in fine chemical production or breakdown.

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Further, the aforementioned enzymes which participate in aromatic/aliphatic compound modification or degradation, general biocatalysis, inorganic compound metabolism or proteolysis are each themselves fine chemicals, desirable for their activity in various *in vitro* industrial applications. By altering the number of copies of the gene for one or more of these enzymes in *C. glutumicum* it may be possible to increase the number of these proteins produced by the cell, thereby increasing the potential yield or efficiency of production of these proteins from large-scale *C. glutumicum* or related bacterial cultures.

The alteration of an HA protein of the invention may also indirectly affect the yield, production, and/or efficiency of production of a fine chemical from a C. glutamicum strain incorporating such an altered protein. For example, by modulating the activity and/or number of those proteins involved in the construction or rearrangement of the cell wall, it may be possible to modify the structure of the cell wall itself such that the cell is able to better withstand the mechanical and other stresses present during large-scale fermentative culture. Also, large-scale growth of C glutamicum requires significant cell wall production. Modulation of the activity or number of cell wall biosynthetic or degradative enzymes may allow more rapid rates of cell wall biosynthesis, which in turn may permit increased growth rates of this microorganism in culture and thereby increase the number of cells producing the desired fine chemical.

By modifying the HA enzymes of the invention, one may also indirectly impact the yield, production, or efficiency of production of one or more fine chemicals from C. glutamicum. For example, many of the general enzymes in C. glutamicum may have a significant impact on global cellular processes (e.g., regulatory processes) which in turn have a significant effect on fine chemical metabolism. Similarly, proteases, enzymes which modify or degrade possibly toxic aromatic or aliphatic compounds, and enzymes which promote the metabolism of inorganic compounds all serve to increase the viability of C glutamicum. The proteases aid in the selective removal of misfolded or misregulated proteins, such as those that might occur under the relatively stressful environmental conditions encountered during large-scale fermentor culture. By altering these proteins, it may be possible to further enhance this activity and to improve the viability of C glutamicum in culture. The aromatic/aliphatic modification or degradation proteins not only serve to detoxify these waste compounds (which may be

encountered as impurities in culture medium or as waste products from cells themselves), but also to permit the cells to utilize alternate carbon sources if the optimal carbon source is limiting in the culture. By increasing their number and/or activity, the survival of C. glutamicum cells in culture may be enhanced. The inorganic metabolism proteins of the invention supply the cell with inorganic molecules required for all protein and nucleotide (among others) synthesis, and thus are critical for the overall viability of the cell. An increase in the number of viable cells producing one or more desired fine chemicals in large-scale culture should result in a concomitant increase in the yield, production, and/or efficiency of production of the fine chemical in the culture.

The isolated nucleic acid sequences of the invention are contained within the genome of a Corynebacterium glutamicum strain available through the American Type Culture Collection, given designation ATCC 13032. The nucleotide sequence of the isolated C. glutamicum HA cDNAs and the predicted amino acid sequences of the C glutamicum HA proteins are shown in Appendices A and B, respectively.

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Computational analyses were performed which classified and/or identified these nucleotide sequences as sequences which encode proteins that participate in Cgluramicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or that have a C. glutamicum enzymatic or proteolytic activity.

The present invention also pertains to proteins which have an amino acid sequence which is substantially homologous to an amino acid sequence of Appendix B. As used herein, a protein which has an amino acid sequence which is substantially homologous to a selected amino acid sequence is least about 50% homologous to the selected amino acid sequence, e.g., the entire selected amino acid sequence. A protein 25 which has an amino acid sequence which is substantially homologous to a selected amino acid sequence can also be least about 50-60%, preferably at least about 60-70%, and more preferably at least about 70-80%, 80-90%, or 90-95%, and most preferably at least about 96%, 97%, 98%, 99% or more homologous to the selected amino acid sequence.

The HA protein or a biologically active portion or fragment thereof of the invention can participate in the maintenance of homeostasis in C. glutumicum, or can perform a function involved in the adaptation of this microorganism to different environmental conditions, or have one or more of the activities set forth in Table 1.

Various aspects of the invention are described in further detail in the following subsections.

## A Isolated Nucleic Acid Molecules

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One aspect of the invention pertains to isolated nucleic acid molecules that encode HA polypeptides or biologically active portions thereof, as well as nucleic acid fragments sufficient for use as hybridization probes or primers for the identification or amplification of HA-encoding nucleic acid (e.g., HA DNA). As used herein, the term "nucleic acid molecule" is intended to include DNA molecules (e.g., cDNA or genomic DNA) and RNA molecules (e.g., mRNA) and analogs of the DNA or RNA generated using nucleotide analogs. This term also encompasses untranslated sequence located at both the 3' and 5' ends of the coding region of the gene: at least about 100 nucleotides of sequence upstream from the 5' end of the coding region and at least about 20 nucleotides of sequence downstream from the 3'end of the coding region of the gene. The nucleic acid molecule can be single-stranded or double-stranded, but preferably is double-stranded DNA. An "isolated" nucleic acid molecule is one which is separated from other nucleic acid molecules which are present in the natural source of the nucleic acid. Preferably, an "isolated" nucleic acid is free of sequences which naturally flank the nucleic acid (i.e., sequences located at the 5' and 3' ends of the nucleic acid) in the genomic DNA of the organism from which the nucleic acid is derived. For example, in various embodiments, the isolated HA nucleic acid molecule can contain less than about 5 kb, 4kb, 3kb, 2kb, 1 kb, 0.5 kb or 0.1 kb of nucleotide sequences which naturally flank the nucleic acid molecule in genomic DNA of the cell from which the nucleic acid is derived (e.g. a C. glutamicum cell). Moreover, an "isolated" nucleic acid molecule, such as a cDNA molecule, can be substantially free of other cellular material, or culture medium when produced by recombinant techniques, or chemical precursors or other chemicals when chemically synthesized.

A nucleic acid molecule of the present invention, e.g., a nucleic acid molecule having a nucleotide sequence of Appendix A, or a portion thereof, can be isolated using standard molecular biology techniques and the sequence information provided herein. For example, a C. glutamicum HA cDNA can be isolated from a C. glutamicum library using all or portion of one of the sequences of Appendix A as a hybridization probe and standard hybridization techniques (e.g., as described in Sambrook, J., Fritsh, E. F., and Maniatis, T. Molecular Cloning: A Laboratory Manual. 2nd. ed. Cold Spring Harbor Laboratory. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989). Moreover, a nucleic acid molecule encompassing all or a portion of one of the sequences of Appendix A can be isolated by the polymerase chain reaction using oligonucleotide primers designed based upon this sequence (e.g., a nucleic acid molecule encompassing all or a portion of one of the sequences of Appendix A can be isolated by the polymerase chain reaction using oligonucleotide primers designed based upon this same sequence of

Appendix A). For example, mRNA can be isolated from normal endothelial cells (e.g., by the guanidinium-thiocyanate extraction procedure of Chirgwin et al. (1979) Biochemistry 18: 5294-5299) and cDNA can be prepared using reverse transcriptase (e.g., Moloney MLV reverse transcriptase, available from Gibco/BRL, Bethesda, MD; or AMV reverse transcriptase, available from Seikagaku America, Inc., St. Petersburg, FL). Synthetic oligonucleotide primers for polymerase chain reaction amplification can be designed based upon one of the nucleotide sequences shown in Appendix A. A nucleic acid of the invention can be amplified using cDNA or, alternatively, genomic DNA, as a template and appropriate oligonucleotide primers according to standard PCR amplification techniques. The nucleic acid so amplified can be cloned into an appropriate vector and characterized by DNA sequence analysis. Furthermore, oligonucleotides corresponding to an HA nucleotide sequence can be prepared by standard synthetic techniques, e.g., using an automated DNA synthesizer.

In a preferred embodiment, an isolated nucleic acid molecule of the invention comprises one of the nucleotide sequences shown in Appendix A. The sequences of Appendix A correspond to the Corynebacterium glutamicum HA cDNAs of the invention. This cDNA comprises sequences encoding HA proteins (i.e., "the coding region", indicated in each sequence in Appendix A), as well as 5' untranslated sequences and 3' untranslated sequences, also indicated in Appendix A. Alternatively, the nucleic acid molecule can comprise only the coding region of any of the sequences in Appendix

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For the purposes of this application, it will be understood that each of the sequences set forth in Appendix A has an identifying RXA number having the designation "RXA" followed by 5 digits (i.e., RXA00009). Each of these sequences comprises up to three parts: a 5' upstream region, a coding region, and a downstream region. Each of these three regions is identified by the same RXA designation to eliminate confusion. The recitation "one of the sequences in Appendix A", then, refers to any of the sequences in Appendix A, which may be distinguished by their differing RXA designations. The coding region of each of these sequences is translated into a 30 corresponding amino acid sequence, which is set forth in Appendix B. The sequences of Appendix B are identified by the same RXA designations as Appendix A, such that they can be readily correlated. For example, the amino acid sequence in Appendix B designated RXA00009 is a translation of the coding region of the nucleotide sequence of nucleic acid molecule RXA00009 in Appendix A.

In one embodiment, the nucleic acid molecules of the present invention are not **35** . intended to include those compiled in Table 2. In the case of the dapD gene, a sequence for this gene was published in Wehrmann, A., et al. (1998) J. Bucteriol. 180(12): 3159-

3165. However, the sequence obtained by the inventors of the present application is significantly longer than the published version. It is believed that the published version relied on an incorrect start codon, and thus represents only a fragment of the actual coding region.

In another preferred embodiment, an isolated nucleic acid molecule of the invention comprises a nucleic acid molecule which is a complement of one of the nucleotide sequences shown in Appendix A, or a portion thereof. A nucleic acid molecule which is complementary to one of the nucleotide sequences shown in Appendix A is one which is sufficiently complementary to one of the nucleotide sequences shown in Appendix A such that it can hybridize to one of the nucleotide sequences shown in Appendix A, thereby forming a stable duplex.

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In still another preferred embodiment, an isolated nucleic acid molecule of the invention comprises a nucleotide sequence which is at least about 50-60%, preferably at least about 60-70%, more preferably at least about 70-80%, 80-90%, or 90-95%, and even more preferably at least about 95%, 96%, 97%, 98%, 99% or more homologous to a nucleotide sequence shown in Appendix A, or a portion thereof. In an additional preferred embodiment, an isolated nucleic acid molecule of the invention comprises a nucleotide sequence which hybridizes, e.g., hybridizes under stringent conditions, to one of the nucleotide sequences shown in Appendix A, or a portion thereof.

Moreover, the nucleic acid molecule of the invention can comprise only a portion of the coding region of one of the sequences in Appendix A, for example a fragment which can be used as a probe or primer or a fragment encoding a biologically active portion of an HA protein. The nucleonide sequences determined from the cloning of the HA genes from C. glutamicum allows for the generation of probes and primers designed for use in identifying and/or cloning HA homologues in other cell types and organisms, as well as HA homologues from other Corynebacteria or related species. The probe/primer typically comprises substantially purified oligonucleotide. The oligonucleotide typically comprises a region of nucleotide sequence that hybridizes under stringent conditions to at least about 12, preferably about 25, more preferably about 40, 50 or 75 consecutive nucleotides of a sense strand of one of the sequences set forth in Appendix A, an anti-sense sequence of one of the sequences set forth in Appendix A, or naturally occurring mutants thereof. Primers based on a nucleotide sequence of Appendix A can be used in PCR reactions to clone HA homologues. Probes based on the HA nucleotide sequences can be used to detect transcripts or genomic sequences encoding the same or homologous proteins. In preferred embodiments, the probe further comprises a label group attached thereto, e.g. the label group can be a radioisotope, a fluorescent compound, an enzyme, or an enzyme cofactor. Such probes can be used as a part of a diagnostic test kit for identifying cells which misexpress an HA protein, such as by measuring a level of an HA-encoding nucleic acid in a sample of cells, e.g., detecting HA mRNA levels or determining whether a genomic HA gene has been mutated or deleted.

In one embodiment, the nucleic acid molecule of the invention encodes a protein or portion thereof which includes an amino acid sequence which is sufficiently homologous to an amino acid sequence of Appendix B such that the protein or portion thereof maintains the ability to participate in the maintenance of homeostasis in C' glutumicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions. As used herein, the language "sufficiently homologous" refers to proteins or portions thereof which have amino acid sequences which include a minimum number of identical or equivalent (e.g., an amino acid residue which has a similar side chain as an amino acid residue in one of the sequences of Appendix B) amino acid residues to an amino acid sequence of Appendix B such that the protein or portion thereof is able to participate in the maintenance of homeostasis in C. glutamicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions. Proteins involved in C. glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or aliphatic compounds, or that have a C. glutamicum enzymatic or proteolytic activity, as described herein, may play a role in the production and secretion of one or more fine chemicals. Examples of such activities are also described herein. Thus, "the function of an HA protein" contributes either directly or indirectly to the yield, production, and/or efficiency of production of one or more fine chemicals. Examples of HA protein activities are set forth in Table 1.

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In another embodiment, the protein is at least about 50-60%, preferably at least about 60-70%, and more preferably at least about 70-80%, 80-90%, 90-95%, and most preferably at least about 96%, 97%, 98%, 99% or more homologous to an entire amino acid sequence of Appendix B.

Portions of proteins encoded by the HA nucleic acid molecules of the invention are preferably biologically active portions of one of the HA proteins. As used herein, the term "biologically active portion of an HA protein" is intended to include a portion, e.g., a domain/motif, of an HA protein that can participate in the maintenance of homeostasis in C. glutamicum, or that can perform a function involved in the adaptation of this microorganism to different environmental conditions, or has an activity as set forth in Table 1. To determine whether an HA protein or a biologically active portion thereof can participate in C glutamicum cell wall biosynthesis or rearrangements, metabolism of inorganic compounds, modification or degradation of aromatic or

aliphatic compounds, or has a C glutamicum enzymatic or proteolytic activity, an assay of enzymatic activity may be performed. Such assay methods are well known to those skilled in the art, as detailed in Example 8 of the Exemplification.

Additional nucleic acid fragments encoding biologically active portions of an HA protein can be prepared by isolating a portion of one of the sequences in Appendix B, expressing the encoded portion of the HA protein or peptide (e.g., by recombinant expression *in vitro*) and assessing the activity of the encoded portion of the HA protein or peptide.

The invention further encompasses nucleic acid molecules that differ from one of the nucleotide sequences shown in Appendix A (and portions thereof) due to degeneracy of the genetic code and thus encode the same HA protein as that encoded by the nucleotide sequences shown in Appendix A. In another embodiment, an isolated nucleic acid molecule of the invention has a nucleotide sequence encoding a protein having an amino acid sequence shown in Appendix B. In a still further embodiment, the nucleic acid molecule of the invention encodes a full length C. glutamicum protein which is substantially homologous to an amino acid sequence of Appendix B (encoded by an open reading frame shown in Appendix A).

In addition to the C glutamicum HA nucleotide sequences shown in Appendix A, it will be appreciated by those skilled in the art that DNA sequence polymorphisms that lead to changes in the amino acid sequences of HA proteins may exist within a population (e.g., the C. glutamicum population). Such genetic polymorphism in the HA gene may exist among individuals within a population due to natural variation. As used herein, the terms "gene" and "recombinant gene" refer to nucleic acid molecules comprising an open reading frame encoding an HA protein, preferably a C. glutamicum HA protein. Such natural variations can typically result in 1-5% variance in the nucleotide sequence of the HA gene. Any and all such nucleotide variations and resulting amino acid polymorphisms in HA that are the result of natural variation and that do not alter the functional activity of HA proteins are intended to be within the scope of the invention.

Nucleic acid molecules corresponding to natural variants and non-C. glutamicum homologues of the C glutamicum HA cDNA of the invention can be isolated based on their homology to the C glutamicum HA nucleic acid disclosed herein using the C glutamicum cDNA; or a portion thereof, as a hybridization probe according to standard hybridization techniques under stringent hybridization conditions. Accordingly, in another embodiment, an isolated nucleic acid molecule of the invention is at least 15 nucleotides in length and hybridizes under stringent conditions to the nucleic acid molecule comprising a nucleotide sequence of Appendix A. In other embodiments, the



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nucleic acid is at least 30, 50, 100, 250 or more nucleotides in length. As used herein, the term "hybridizes under stringent conditions" is intended to describe conditions for hybridization and washing under which nucleotide sequences at least 60% homologous to each other typically remain hybridized to each other. Preferably, the conditions are such that sequences at least about 65%, more preferably at least about 70%, and even more preferably at least about 75% or more homologous to each other typically remain hybridized to each other. Such stringent conditions are known to those skilled in the art and can be found in Current Protocols in Molecular Biology, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. A preferred, non-limiting example of stringent hybridization conditions are hybridization in 6X sodium chloride/sodium citrate (SSC) at about 45°C, followed by one or more washes in 0.2 X SSC, 0.1% SDS at 50-65°C. Preferably, an isolated nucleic acid molecule of the invention that hybridizes under stringent conditions to a sequence of Appendix A corresponds to a naturally-occurring nucleic acid molecule. As used herein, a "naturally-occurring" nucleic acid molecule refers to an RNA or DNA molecule having a nucleotide sequence that occurs in nature (e.g., encodes a natural protein). In one embodiment, the nucleic acid encodes a natural C glutumicum HA protein.

In addition to naturally-occurring variants of the HA sequence that may exist in the population, the skilled artisan will further appreciate that changes can be introduced by mutation into a nucleotide sequence of Appendix A, thereby leading to changes in the amino acid sequence of the encoded HA protein, without altering the functional ability of the HA protein. For example, nucleotide substitutions leading to amino acid substitutions at "non-essential" amino acid residues can be made in a sequence of Appendix A. A, "non-essential" amino acid residue is a residue that can be altered from the wild-type sequence of one of the HA proteins (Appendix B) without altering the activity of said HA protein, whereas an "essential" amino acid residue is required for HA protein activity. Other amino acid residues, however, (e.g., those that are not conserved or only semi-conserved in the domain having HA activity) may not be essential for activity and thus are likely to be amenable to alteration without altering HA 30 activity

Accordingly, another aspect of the invention pertains to nucleic acid molecules encoding HA proteins that contain changes in amino acid residues that are not essential for HA activity. Such HA proteins differ in amino acid sequence from a sequence contained in Appendix B yet retain at least one of the HA activities described herein. In one embodiment, the isolated nucleic acid molecule comprises a nucleotide sequence encoding a protein, wherein the protein comprises an amino acid sequence at least about 50% homologous to an amino acid sequence of Appendix B and is capable of



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participating in the maintenance of homeostasis in C glutamicum, or of performing a function involved in the adaptation of this microorganism to different environmental conditions, or has one or more of the activities set forth in Table 1. Preferably, the protein encoded by the nucleic acid molecule is at least about 50-60% homologous to one of the sequences in Appendix B, more preferably at least about 60-70% homologous to one of the sequences in Appendix B, even more preferably at least about 70-80%, 80-90%, 90-95% homologous to one of the sequences in Appendix B, and most preferably at least about 96%, 97%, 98%, or 99% homologous to one of the sequences in Appendix B.

To determine the percent homology of two amino acid sequences (e.g., one of the sequences of Appendix B and a mutant form thereof) or of two nucleic acids, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in the sequence of one protein or nucleic acid for optimal alignment with the other protein or nucleic acid). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in one sequence (e.g., one of the sequences of Appendix B) is occupied by the same amino acid residue or nucleotide as the corresponding position in the other sequence (e.g., a mutant form of the sequence selected from Appendix B), then the molecules are homologous at that position (i.e., as used herein amino acid or nucleic acid "homology" is equivalent to amino acid or nucleic acid "identity"). The percent homology between the two sequences is a function of the number of identical positions shared by the sequences (i.e., % homology = # of identical positions/total # of positions x 100).

An isolated nucleic acid molecule encoding an HA protein homologous to a protein sequence of Appendix B can be created by introducing one or more nucleotide substitutions, additions or deletions into a nucleotide sequence of Appendix A such that one or more amino acid substitutions, additions or deletions are introduced into the encoded protein. Mutations can be introduced into one of the sequences of Appendix A by standard techniques, such as site-directed mutagenesis and PCR-mediated mutagenesis. Preferably, conservative amino acid substitutions are made at one or more predicted non-essential amino acid residues. A "conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a similar side chain. Families of amino acid residues having similar side chains have been defined in the art. These families include amino acids with basic side chains (e.g., lysine, arginine, histidine), acidic side chains (e.g., aspartic acid, glutamic acid), uncharged polar side chains (e.g., glycine, asparagine, glutamine, serine, threonine, tyrosine, cysteine), nonpolar side chains (e.g., alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan), beta-branched side chains (e.g.,

threonine, valine, isoleucine) and aromatic side chains (e.g., tyrosine, phenylalanine, tryptophan, histidine). Thus, a predicted nonessential amino acid residue in an HA protein is preferably replaced with another amino acid residue from the same side chain family. Alternatively, in another embodiment, mutations can be introduced randomly along all or part of an HA coding sequence, such as by saturation mutagenesis, and the resultant mutants can be screened for an HA activity described herein to identify mutants that retain HA activity. Following mutagenesis of one of the sequences of Appendix A, the encoded protein can be expressed recombinantly and the activity of the protein can be determined using, for example, assays described herein (see Example 8 of the Exemplification).

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In addition to the nucleic acid molecules encoding HA proteins described above, another aspect of the invention pertains to isolated nucleic acid molecules which are antisense thereto. An "antisense" nucleic acid comprises a nucleotide sequence which is complementary to a "sense" nucleic acid encoding a protein, e.g., complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. Accordingly, an antisense nucleic acid can hydrogen bond to a sense nucleic acid. The antisense nucleic acid can be complementary to an entire HA coding strand, or to only a portion thereof. In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence encoding an HA protein. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid residues (e.g., the entire coding region of SEQ ID RXA00009 comprises nucleotides 1 to 900). In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence encoding HA. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (i.e., also referred to as 5' and 3' untranslated regions).

Given the coding strand sequences encoding HA disclosed herein (e.g., the sequences set forth in Appendix A), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of HA mRNA, but more preferably is an oligonucleotide which is antisense to only a portion of the coding or noncoding region of HA mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of HA mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis and enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can

be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used. Examples of modified nucleotides which can be used to generate the antisense nucleic acid include 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-10 methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-Dmannosylqueosine, 5-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-15 amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (i.e., RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target 20 nucleic acid of interest, described further in the following subsection).

The antisense nucleic acid molecules of the invention are typically administered to a cell or generated in situ such that they hybridize with or bind to cellular mRNA and/or genomic DNA encoding an HA protein to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule which binds to DNA duplexes, through specific interactions in the major groove of the double helix. The antisense molecule can be modified such that it specifically binds to a receptor or an antigen expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecule to a peptide or an antibody which binds to a cell surface receptor or antigen. The antisense nucleic acid molecule can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of the antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong prokaryotic, viral, or eukaryotic promoter are preferred.

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In yet another embodiment, the antisense nucleic acid molecule of the invention is an  $\alpha$ -anomeric nucleic acid molecule. An  $\alpha$ -anomeric nucleic acid molecule forms

specific double-stranded hybrids with complementary RNA in which, contrary to the usual β-units, the strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids. Res. 15:6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res. 15:6131-6148) or a chimeric RNA-DNA analogue (Inoue et al. (1987) FEBS Lett. 215:327-330).

In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity which are capable of cleaving a single-stranded nucleic acid, such as an mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) Nature 334:585-591)) can be used to catalytically cleave HA mRNA transcripts to thereby inhibit translation of HA mRNA. A ribozyme having specificity for an IIA-encoding nucleic acid can be designed based upon the nucleotide sequence of an HA DNA molecule disclosed herein (i.e., RXA00009 in Appendix A). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is complementary to the nucleotide sequence to be cleaved in an HA-encoding mRNA. See, e.g., Cech et al. U.S. Patent No. 4,987,071 and Cech et al. U.S. Patent No. 5,116,742. Alternatively, HA mRNA can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, e.g., Bartel, D. and Szostak, J.W. (1993) Science 261:1411-1418.

Alternatively, HA gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region of an HA nucleotide sequence (e.g., an HA promoter and/or enhancers) to form triple helical structures that prevent transcription of an HA gene in target cells. See generally, Helene, C. (1991) Anticuncer Drug Des. 6(6):569-84; Helene, C. et al. (1992) Ann N.Y. Acad Sci. 660:27-36; and Maher, L.J. (1992) Bioassays 14(12):807-15.

## B. Recombinant Expression Vectors and Host Cells

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Another aspect of the invention pertains to vectors, preferably expression vectors, containing a nucleic acid encoding an HA protein (or a portion thereof). As used herein, the term "vector" refers to a nucleic acid molecule capable of transporting another nucleic acid to which it has been linked. One type of vector is a "plasmid", which refers to a circular double stranded DNA loop into which additional DNA segments can be ligated. Another type of vector is a viral vector, wherein additional DNA segments can be ligated into the viral genome. Certain vectors are capable of autonomous replication in a host cell into which they are introduced (e.g., bacterial vectors having a bacterial origin of replication and episomal mammalian vectors). Other

vectors (e.g., non-episomal mammalian vectors) are integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome. Moreover, certain vectors are capable of directing the expression of genes to which they are operatively linked. Such vectors are referred to herein as "expression vectors". In general, expression vectors of utility in recombinant DNA techniques are often in the form of plasmids. In the present specification, "plasmid" and "vector" can be used interchangeably as the plasmid is the most commonly used form of vector. However, the invention is intended to include such other forms of expression vectors, such as viral vectors (e.g., replication defective retroviruses, adenoviruses and adenoassociated viruses), which serve equivalent functions.

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The recombinant expression vectors of the invention comprise a nucleic acid of the invention in a form suitable for expression of the nucleic acid in a host cell, which means that the recombinant expression vectors include one or more regulatory sequences, selected on the basis of the host cells to be used for expression, which are operatively linked to the nucleic acid sequence to be expressed. Within a recombinant expression vector, "operably linked" is intended to mean that the nucleotide sequence of interest is linked to the regulatory sequence(s) in a manner which allows for expression of the nucleotide sequence (e.g., in an in vitro transcription/translation system or in a host cell when the vector is introduced into the host cell). The term "regulatory sequence" is intended to include promoters, enhancers and other expression control elements (e.g., polyadenylation signals). Such regulatory sequences are described, for example, in Goeddel; Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, CA (1990). Regulatory sequences include those which direct constitutive expression of a nucleotide sequence in many types of host cell and those which direct expression of the nucleotide sequence only in certain host cells. It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host cell to be transformed, the level of expression of protein desired, etc. The expression vectors of the invention can be introduced into host cells to thereby produce proteins or peptides, including fusion proteins or peptides, encoded by nucleic acids as described herein (e.g., HA proteins, mutant forms of HA proteins, fusion proteins, etc.).

The recombinant expression vectors of the invention can be designed for expression of HA proteins in prokaryotic or eukaryotic cells. For example, HA genes can be expressed in bacterial cells such as C glutamicum, insect cells (using baculovirus expression vectors), yeast and other fungal cells (see Romanos, M.A. et al. (1992) "Foreign gene expression in yeast: a review", Yeast 8: 423-488; van den Hondel, C.A.M.J.J. et al. (1991) "Heterologous gene expression in filamentous fungi" in: More

Gene Manipulations in Fungi, J.W. Bennet & L.L. Lasure, eds., p. 396-428: Academic Press: San Diego; and van den Hondel, C.A.M.J.J. & Punt, P.J. (1991) "Gene transfer systems and vector development for filamentous fungi, in: Applied Molecular Genetics of Fungi, Peberdy, J.F. et al., eds., p. 1-28, Cambridge University Press: Cambridge), algae and multicellular plant cells (see Schmidt, R. and Willmitzer, L. (1988) High efficiency Agrobacterium tumefaciens—mediated transformation of Arabidopsis thatiana leaf and cotyledon explants" Plant Cell Rep.: 583-586), or mammalian cells. Suitable host cells are discussed further in Goeddel, Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, CA (1990). Alternatively, the recombinant expression vector can be transcribed and translated in vitro, for example using T7 promoter regulatory sequences and T7 polymerase.

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Expression of proteins in prokaryotes is most often carried out with vectors containing constitutive or inducible promoters directing the expression of either fusion or non-fusion proteins. Fusion vectors add a number of amino acids to a protein encoded therein, usually to the amino terminus of the recombinant protein but also to the C-terminus or fused within suitable regions in the proteins. Such fusion vectors typically serve three purposes: 1) to increase expression of recombinant protein; 2) to increase the solubility of the recombinant protein; and 3) to aid in the purification of the recombinant protein by acting as a ligand in affinity purification. Often, in fusion expression vectors, a proteolytic cleavage site is introduced at the junction of the fusion moiety and the recombinant protein to enable separation of the recombinant protein from the fusion moiety subsequent to purification of the fusion protein. Such enzymes, and their cognate recognition sequences, include Factor Xa, thrombin and enterokinase.

Typical fusion expression vectors include pGEX (Pharmacia Biotech Inc; Smith, D.B. and Johnson, K.S. (1988) Gene 67:31-40), pMAL (New England Biolabs, Beverly, MA) and pRIT5 (Pharmacia, Piscataway, NJ) which fuse glutathione S-transferase (GST), maltose E binding protein, or protein A, respectively, to the target recombinant protein. In one embodiment, the coding sequence of the HA protein is cloned into a pGEX expression vector to create a vector encoding a fusion protein comprising, from the N-terminus to the C-terminus, GST-thrombin cleavage site-X protein. The fusion protein can be purified by affinity chromatography using glutathione-agarose resin. Recombinant HA protein unfused to GST can be recovered by cleavage of the fusion protein with thrombin.

Examples of suitable inducible non-fusion E. coli expression vectors include pTrc (Amann et al., (1988) Gene 69:301-315) and pET 11d (Studier et al., Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, California (1990) 60-89). Target gene expression from the pTrc vector relies on host

RNA polymerase transcription from a hybrid trp-lac fusion promoter. Target gene expression from the pET 11d vector relies on transcription from a T7 gn10-lac fusion promoter mediated by a coexpressed viral RNA polymerase (T7 gn1). This viral polymerase is supplied by host strains BL21(DE3) or HMS174(DE3) from a resident  $\lambda$  prophage harboring a T7 gn1 gene under the transcriptional control of the lacUV 5 promoter.

One strategy to maximize recombinant protein expression is to express the protein in a host bacteria with an impaired capacity to proteolytically cleave the recombinant protein (Gottesman, S., Gene Expression Technology Methods in Enzymology 185, Academic Press, San Diego, California (1990) 119-128). Another strategy is to alter the nucleic acid sequence of the nucleic acid to be inserted into an expression vector so that the individual codons for each amino acid are those preferentially utilized in the bacterium chosen for expression, such as C. glutamicum (Wada et al. (1992) Nucleic Acids Res. 20: 2111-2118). Such alteration of nucleic acid sequences of the invention can be carried out by standard DNA synthesis techniques.

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In another embodiment, the HA protein expression vector is a yeast expression vector. Examples of vectors for expression in yeast S. cerivisue include pYepSec1 (Baldari, et al., (1987) Embo J. 6:229-234), pMFa (Kurjan and Herskowitz, (1982) Cell 30:933-943), pJRY88 (Schultz et al., (1987) Gene 54:113-123), and pYES2 (Invitrogen Corporation, San Diego, CA). Vectors and methods for the construction of vectors appropriate for use in other fungi, such as the filamentous fungi, include those detailed in: van den Hondel, C.A.M.J.J. & Punt, P.J. (1991) "Gene transfer systems and vector development for filamentous fungi, in: Applied Molecular Genetics of Fungi, J.F. Peberdy, et al., eds., p. 1-28, Cambridge University Press: Cambridge.

Alternatively, the HA proteins of the invention can be expressed in insect cells using baculovirus expression vectors. Baculovirus vectors available for expression of proteins in cultured insect cells (e.g., Sf 9 cells) include the pAc series (Smith et al. (1983) Mol Cell Biol 3:2156-2165) and the pVL series (Lucklow and Summers (1989) Virology 170:31-39).

In another embodiment, the HA proteins of the invention may be expressed in unicellular plant cells (such as algae) or in plant cells from higher plants (e.g., the spermatophytes, such as crop plants). Examples of plant expression vectors include those detailed in: Becker, D., Kemper, E., Schell, J. and Masterson, R. (1992) "New plant binary vectors with selectable markers located proximal to the left border", *Plant Mol. Biol.* 20. 1195-1197; and Bevan, M.W. (1984) "Binary Agrobacterium vectors for plant transformation", *Nucl. Acid. Res.* 12: 8711-8721.

In yet another embodiment, a nucleic acid of the invention is expressed in mammalian cells using a mammalian expression vector. Examples of mammalian expression vectors include pCDM8 (Seed, B. (1987) Nature 329:840) and pMT2PC (Kaufman et al. (1987) EMBO J. 6:187-195). When used in mammalian cells, the expression vector's control functions are often provided by viral regulatory elements. For example, commonly used promoters are derived from polyoma, Adenovirus 2, cytomegalovirus and Simian Virus 40. For other suitable expression systems for both prokaryotic and eukaryotic cells see chapters 16 and 17 of Sambrook, J., Fritsh, E. F., and Maniatis, T. Molecular Cloning. A Laboratory Manual. 2nd, ed., Cold Spring Harbor Laboratory. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989.

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In another embodiment, the recombinant mammalian expression vector is capable of directing expression of the nucleic acid preferentially in a particular cell type (e.g., tissue-specific regulatory elements are used to express the nucleic acid). Tissuespecific regulatory elements are known in the art. Non-limiting examples of suitable tissue-specific promoters include the albumin promoter (liver-specific; Pinkert et al. (1987) Genes Dev. 1:268-277), lymphoid-specific promoters (Calame and Eaton (1988) Adv. Immunol 43:235-275), in particular promoters of T cell receptors (Winoto and Baltimore (1989) EMBO J. 8:729-733) and immunoglobulins (Banerji et al. (1983) Cell 33:729-740; Queen and Baltimore (1983) Cell 33:741-748), neuron-specific promoters (e.g., the neurofilament promoter; Byrne and Ruddle (1989) PNAS 86:5473-5477), pancreas-specific promoters (Edlund et al. (1985) Science 230:912-916), and mammary gland-specific promoters (e.g., milk whey promoter; U.S. Patent No. 4,873,316 and European Application Publication No. 264,166). Developmentally-regulated promoters are also encompassed, for example the murine hox promoters (Kessel and Gruss (1990) Science 249:374-379) and the  $\alpha$ -fetoprotein promoter (Campes and Tilghman (1989) Genes Dev. 3:537-546).

The invention further provides a recombinant expression vector comprising a DNA molecule of the invention cloned into the expression vector in an antisense orientation. That is, the DNA molecule is operatively linked to a regulatory sequence in a manner which allows for expression (by transcription of the DNA molecule) of an RNA molecule which is antisense to HA mRNA. Regulatory sequences operatively linked to a nucleic acid cloned in the antisense orientation can be chosen which direct the continuous expression of the antisense RNA molecule in a variety of cell types, for instance viral promoters and/or enhancers, or regulatory sequences can be chosen which direct constitutive, tissue specific or cell type specific expression of antisense RNA. The antisense expression vector can be in the form of a recombinant plasmid, phagemid

or attenuated virus in which antisense nucleic acids are produced under the control of a high efficiency regulatory region, the activity of which can be determined by the cell type into which the vector is introduced. For a discussion of the regulation of gene expression using antisense genes see Weintraub, H. et al., Antisense RNA as a molecular tool for genetic analysis, Reviews - Trends in Genetics, Vol. 1(1) (1986).

Another aspect of the invention pertains to host cells into which a recombinant expression vector of the invention has been introduced. The terms "host cell" and "recombinant host cell" are used interchangeably herein. It is understood that such terms refer not only to the particular subject cell but to the progeny or potential progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not, in fact, be identical to the parent cell, but are still included within the scope of the term as used herein.

A host cell can be any prokaryotic or eukaryotic cell. For example, an HA protein can be expressed in bacterial cells such as *C. glutamicum*, insect cells, yeast or mammalian cells (such as Chinese hamster ovary cells (CHO) or COS cells). Other suitable host cells are known to those skilled in the art. Microorganisms related to *Corynebacterium glutamicum* which may be conveniently used as host cells for the nucleic acid and protein molecules of the invention are set forth in Table 3.

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Vector DNA can be introduced into prokaryotic or eukaryotic cells via conventional transformation or transfection techniques. As used herein, the terms "transformation" and "transfection", "conjugation" and "transduction" are intended to refer to a variety of art-recognized techniques for introducing foreign nucleic acid (e.g., DNA) into a host cell, including calcium phosphate or calcium chloride co-precipitation, DEAE-dextran-mediated transfection, lipofection, natural competence, chemical-mediated transfer, or electroporation. Suitable methods for transforming or transfecting host cells can be found in Sambrook, et al. (Molecular Cloning: A Laboratory Manual. 2nd, Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989), and other laboratory manuals.

For stable transfection of mammalian cells, it is known that, depending upon the expression vector and transfection technique used, only a small fraction of cells may integrate the foreign DNA into their genome. In order to identify and select these integrants, a gene that encodes a selectable marker (e.g., resistance to antibiotics) is generally introduced into the host cells along with the gene of interest. Preferred selectable markers include those which confer resistance to drugs, such as G418, hygromycin and methotrexate. Nucleic acid encoding a selectable marker can be introduced into a host cell on the same vector as that encoding an HA protein or can be

introduced on a separate vector. Cells stably transfected with the introduced nucleic acid can be identified by, for example, drug selection (e.g., cells that have incorporated the selectable marker gene will survive, while the other cells die).

To create a homologous recombinant microorganism, a vector is prepared which contains at least a portion of an HA gene into which a deletion, addition or substitution has been introduced to thereby alter, e.g., functionally disrupt, the HA gene. Preferably, this HA gene is a Corynebacterium glutamicum HA gene, but it can be a homologue from a related bacterium or even from a mammalian, yeast, or insect source. In a preferred embodiment, the vector is designed such that, upon homologous 10 recombination, the endogenous HA gene is functionally disrupted (i.e., no longer encodes a functional protein; also referred to as a "knock out" vector). Alternatively, the vector can be designed such that, upon homologous recombination, the endogenous HA gene is mutated or otherwise altered but still encodes functional protein (e.g., the upstream regulatory region can be altered to thereby alter the expression of the endogenous HA protein). In the homologous recombination vector, the altered portion of the HA gene is flanked at its 5' and 3' ends by additional nucleic acid of the HA gene to allow for homologous recombination to occur between the exogenous HA gene carried by the vector and an endogenous HA gene in a microorganism. The additional flanking HA nucleic acid is of sufficient length for successful homologous 20 recombination with the endogenous gene. Typically, several kilobases of flanking DNA (both at the 5' and 3' ends) are included in the vector (see e.g., Thomas, K.R., and Capecchi, M.R. (1987) Cell 51: 503 for a description of homologous recombination vectors). The vector is introduced into a microorganism (e.g., by electroporation) and cells in which the introduced HA gene has homologously recombined with the endogenous HA gene are selected, using art-known techniques. 25

In another embodiment, recombinant microorganisms can be produced which contain selected systems which allow for regulated expression of the introduced gene. For example, inclusion of an HA gene on a vector placing it under control of the lac operon permits expression of the HA gene only in the presence of IPTG. Such regulatory systems are well known in the art.

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A host cell of the invention, such as a prokaryotic or eukaryotic host cell in culture, can be used to produce (i.e., express) an HA protein. Accordingly, the invention further provides methods for producing HA proteins using the host cells of the invention. In one embodiment, the method comprises culturing the host cell of invention (into which a recombinant expression vector encoding an HA protein has been introduced, or into which genome has been introduced a gene encoding a wild-type or altered HA protein) in a suitable medium until HA protein is produced. In another

embodiment, the method further comprises isolating HA proteins from the medium or the host cell.

### C. Isolated HA Proteins .

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Another aspect of the invention pertains to isolated HA proteins, and biologically . 5 active portions thereof. An "isolated" or "purified" protein or biologically active portion thereof is substantially free of cellular material when produced by recombinant DNA techniques, or chemical precursors or other chemicals when chemically synthesized. The language "substantially free of cellular material" includes preparations of HA protein in which the protein is separated from cellular components of the cells in which it is naturally or recombinantly produced. In one embodiment, the language "substantially free of cellular material" includes preparations of HA protein having less than about 30% (by dry weight) of non-HA protein (also referred to herein as a "contaminating protein"), more preferably less than about 20% of non-HA protein, still more preferably less than about 10% of non-HA protein, and most preferably less than about 5% non-HA protein. When the HA protein or biologically active portion thereof is recombinantly produced, it is also preferably substantially free of culture medium, i.e., culture medium represents less than about 20%, more preferably less than about 10%, and most preferably less than about 5% of the volume of the protein preparation. The language "substantially free of chemical precursors or other chemicals" includes preparations of HA projein in which the protein is separated from chemical précursors or other chemicals which are involved in the synthesis of the protein. In one embodiment, the language "substantially free of chemical precursors or other chemicals" includes preparations of HA protein having less than about 30% (by dry weight) of chemical precursors or non-HA chemicals, more preferably less than about 20% chemical precursors or non-HA chemicals, still more preferably less than about 10% chemical precursors or non-HA chemicals, and most preferably less than about 5% chemical precursors or non-HA chemicals. In preferred embodiments, isolated proteins or biologically active portions thereof lack contaminating proteins from the same organism from which the HA protein is derived. Typically, such proteins are produced by recombinant expression of, for example, a C glutamicum HA protein in a microorganism such as C. glutamicum.

An isolated HA protein or a portion thereof of the invention can participate in the repair or recombination of DNA, in the transposition of genetic material, in gene expression (i.e., the processes of transcription or translation), in protein folding, or in protein secretion in Corynebacterium glutamicum, or has one or more of the activities set forth in Table 1. In preferred embodiments, the protein or portion thereof comprises

an amino acid sequence which is sufficiently homologous to an amino acid sequence of Appendix B such that the protein or portion thereof maintains the ability to participate in the maintenance of homeostasis in C glutamicum, or to perform a function involved in the adaptation of this microorganism to different environmental conditions. The portion 5 of the protein is preferably a biologically active portion as described herein. In another preferred embodiment, an HA protein of the invention has an amino acid sequence shown in Appendix B. In yet another preferred embodiment, the HA protein has an amino acid sequence which is encoded by a nucleotide sequence which hybridizes, e.g., hybridizes under stringent conditions, to a nucleotide sequence of Appendix A. In still another preferred embodiment, the HA protein has an amino acid sequence which is encoded by a nucleotide sequence that is at least about 50-60%, preferably at least about 60-70%, more preferably at least about 70-80%, 80-90%, 90-95%, and even more preferably at least about 96%, 97%, 98%, 99% or more homologous to one of the amino acid sequences of Appendix B. The preferred HA proteins of the present invention also preferably possess at least one of the HA activities described herein. For example, a preferred HA protein of the present invention includes an amino acid sequence encoded by a nucleotide sequence which hybridizes, e.g., hybridizes under stringent conditions, to a nucleotide sequence of Appendix A, and which can participate in the maintenance of homeostasis in C glutamicum, or can perform a function involved in the adaptation of this microorganism to different environmental conditions, or which has one or more of the activities set forth in Table 1.

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In other embodiments, the HA protein is substantially homologous to an amino acid sequence of Appendix B and retains the functional activity of the protein of one of the sequences of Appendix B yet differs in amino acid sequence due to natural variation or mutagenesis, as described in detail in subsection I above. Accordingly, in another embodiment, the HA protein is a protein which comprises an amino acid sequence which is at least about 50-60%, preferably at least about 60-70%, and more preferably at least about 70-80, 80-90, 90-95%, and most preferably at least about 96%, 97%, 98%, 99% or more homologous to an entire amino acid sequence of Appendix B and which has at least one of the HA activities described herein. In another embodiment, the invention pertains to a full length C glutamicum protein which is substantially homologous to an entire amino acid sequence of Appendix B.

Biologically active portions of an HA protein include peptides comprising amino acid sequences derived from the amino acid sequence of an HA protein, e.g., the an amino acid sequence shown in Appendix B or the amino acid sequence of a protein homologous to an HA protein, which include fewer amino acids than a full length HA protein or the full length protein which is homologous to an HA protein, and exhibit at

least one activity of an HA protein. Typically, biologically active portions (peptides, e.g., peptides which are, for example, 5, 10, 15, 20, 30, 35, 36, 37, 38, 39, 40, 50, 100 or more amino acids in length) comprise a domain or motif with at least one activity of an HA protein. Moreover, other biologically active portions, in which other regions of the protein are deleted, can be prepared by recombinant techniques and evaluated for one or more of the activities described herein. Preferably, the biologically active portions of an HA protein include one or more selected domains/motifs or portions thereof having biological activity.

SES proteins are preferably produced by recombinant DNA techniques. For example, a nucleic acid molecule encoding the protein is cloned into an expression vector (as described above), the expression vector is introduced into a host cell (as described above) and the HA protein is expressed in the host cell. The HA protein can then be isolated from the cells by an appropriate purification scheme using standard protein purification techniques. Alternative to recombinant expression, an HA protein, polypeptide, or peptide can be synthesized chemically using standard peptide synthesis techniques. Moreover, native HA protein can be isolated from cells (e.g., endothelial cells), for example using an anti-HA antibody, which can be produced by standard techniques utilizing an HA protein or fragment thereof of this invention.

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The invention also provides HA chimeric or fusion proteins. As used herein, an HA "chimeric protein" or "fusion protein" comprises an HA polypeptide operatively linked to a non-HA polypeptide. An "HA polypeptide" refers to a polypeptide having an amino acid sequence corresponding to an HA protein, whereas a "non-HA polypeptide" refers to a polypeptide having an amino acid sequence corresponding to a protein which is not substantially homologous to the HA protein, e.g., a protein which is different from the HA protein and which is derived from the same or a different organism. Within the fusion protein, the term "operatively linked" is intended to indicate that the HA polypeptide and the non-HA polypeptide are fused in-frame to each other. The non-HA polypeptide can be fused to the N-terminus or C-terminus of the HA polypeptide. For example, in one embodiment the fusion protein is a GST-HA fusion protein in which the HA sequences are fused to the C-terminus of the GST sequences. Such fusion proteins can facilitate the purification of recombinant HA proteins. In another embodiment, the fusion protein is an HA protein containing a heterologous signal sequence at its Nterminus. In certain host cells (e.g., mammalian host cells), expression and/or secretion of an HA protein can be increased through use of a heterologous signal sequence.

Preferably, an HA chimeric or fusion protein of the invention is produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with

conventional techniques, for example by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers which give rise to complementary overhangs between two consecutive gene fragments which can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for example, Current Protocols in Molecular Biology, eds. Ausubel et al. John Wiley & Sons: 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). An HAencoding nucleic acid can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the HA protein.

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Homologues of the HA protein can be generated by mutagenesis, e.g., discrete point mutation or truncation of the HA protein. As used herein, the term "homologue" refers to a variant form of the HA protein which acts as an agonist or antagonist of the activity of the HA protein. An agonist of the HA protein can retain substantially the same, or a subset, of the biological activities of the HA protein. An antagonist of the HA protein can inhibit one or more of the activities of the naturally occurring form of the HA protein, by, for example, competitively binding to a downstream or upstream member of a biochemical cascade which includes the HA protein, by binding to a target molecule with which the HA protein interacts, such that no functional interaction is possible, or by binding directly to the HA protein and inhibiting its normal activity.

In an alternative embodiment, homologues of the HA protein can be identified by screening combinatorial libraries of mutants, e.g., truncation mutants, of the HA protein for HA protein agonist or antagonist activity. In one embodiment, a variegated library of HA variants is generated by combinatorial mutagenesis at the nucleic acid level and is encoded by a variegated gene library. A variegated library of HA variants can be produced by, for example, enzymatically ligating a mixture of synthetic oligonucleotides into gene sequences such that a degenerate set of potential HA sequences is expressible as individual polypeptides, or alternatively, as a set of larger fusion proteins (e.g., for phage display) containing the set of HA sequences therein. There are a variety of methods which can be used to produce libraries of potential HA homologues from a degenerate oligonucleotide sequence. Chemical synthesis of a degenerate gene sequence can be performed in an automatic DNA synthesizer, and the synthetic gene then ligated into an appropriate expression vector. Use of a degenerate set of genes allows for the provision, in one mixture, of all of the sequences encoding

the desired set of potential HA sequences. Methods for synthesizing degenerate oligonucleotides are known in the art (see, e.g., Narang, S.A. (1983) Tetrahedron 39:3; Itakura et al. (1984) Annu. Rev. Biochem. 53:323: Itakura et al. (1984) Science 198:1056; Ike et al. (1983) Nucleic Acid Res. 11:477.

In addition, libraries of fragments of the HA protein coding can be used to generate a variegated population of HA fragments for screening and subsequent selection of homologues of an HA protein. In one embodiment, a library of coding sequence fragments can be generated by treating a double stranded PCR fragment of an HA coding sequence with a nuclease under conditions wherein nicking occurs only about once per molecule, denaturing the double stranded DNA, renaturing the DNA to form double stranded DNA which can include sense/antisense pairs from different nicked products, removing single stranded portions from reformed duplexes by treatment with S1 nuclease, and ligating the resulting fragment library into an expression vector. By this method, an expression library can be derived which encodes N-terminal. C-terminal and internal fragments of various sizes of the HA protein.

Several techniques are known in the art for screening gene products of combinatorial libraries made by point mutations or truncation, and for screening cDNA libraries for gene products having a selected property. Such techniques are adaptable for rapid screening of the gene libraries generated by the combinatorial mutagenesis of HA homologues. The most widely used techniques, which are amenable to high through-put analysis, for screening large gene libraries typically include cloning the gene library into replicable expression vectors, transforming appropriate cells with the resulting library of vectors, and expressing the combinatorial genes under conditions in which detection of a desired activity facilitates isolation of the vector encoding the gene whose product was detected. Recursive ensemble mutagenesis (REM), a new technique which enhances the frequency of functional mutants in the libraries, can be used in combination with the screening assays to identify HA homologues (Arkin and Yourvan (1992) PNAS 89:7811-7815; Delgrave et al. (1993) Protein Engineering 6(3):327-331).

In another embodiment, cell based assays can be exploited to analyze a variegated HA library, using methods well known in the art.

### D Uses and Methods of the Invention

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The nucleic acid molecules, proteins, protein homologues, fusion proteins, primers, vectors, and host cells described herein can be used in one or more of the following methods: identification of C. glutamicum and related organisms; mapping of genomes of organisms related to C. glutamicum: identification and localization of C. glutamicum sequences of interest; evolutionary studies: determination of HA protein

regions required for function; modulation of an HA protein activity; modulation of the metabolism of one or more inorganic compounds; modulation of the modification or degradation of one or more aromatic or aliphatic compounds; modulation of cell wall synthesis or rearrangements; modulation of enzyme activity or proteolysis; and modulation of cellular production of a desired compound, such as a fine chemical.

The HA nucleic acid molecules of the invention have a variety of uses. First, they may be used to identify an organism as being Corynebacterium glutamicum or a close relative thereof. Also, they may be used to identify the presence of C glutamicum or a relative thereof in a mixed population of microorganisms. The invention provides the nucleic acid sequences of a number of C. glutamicum genes; by probing the extracted genomic DNA of a culture of a unique or mixed population of microorganisms under stringent conditions with a probe spanning a region of a C glutamicum gene which is unique to this organism, one can ascertain whether this organism is present. Although Corynebacterium glutamicum itself is nonpathogenic, it is related to pathogenic species, such as Corynebacterium diphtheriae. Detection of such organisms is of significant clinical relevance.

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Further, the nucleic acid and protein molecules of the invention may serve as markers for specific regions of the genome. This has utility not only in the mapping of the genome, but also for functional studies of *C. glutamicum* proteins. For example, to identify the region of the genome to which a particular *C. glutamicum* DNA-binding protein binds, the *C. glutamicum* genome could be digested, and the fragments incubated with the DNA-binding protein. Those which bind the protein may be additionally probed with the nucleic acid molecules of the invention, preferably with readily detectable labels; binding of such a nucleic acid molecule to the genome fragment enables the localization of the fragment to the genome map of *C. glutamicum*, and, when performed multiple times with different enzymes, facilitates a rapid determination of the nucleic acid sequence to which the protein binds. Further, the nucleic acid molecules of the invention may be sufficiently homologous to the sequences of related species such that these nucleic acid molecules may serve as markers for the construction of a genomic map in related bacteria, such as *Brevibacterium lactofermentum*.

The HA nucleic acid molecules of the invention are also useful for evolutionary and protein structural studies. The processes involved in adaptation and the maintenance of homeostasis in which the molecules of the invention participate are utilized by a wide variety of species; by comparing the sequences of the nucleic acid molecules of the present invention to those encoding similar enzymes from other organisms, the evolutionary relatedness of the organisms can be assessed. Similarly, such a comparison permits an assessment of which regions of the sequence are

conserved and which are not, which may aid in determining those regions of the protein which are essential for the functioning of the enzyme. This type of determination is of value for protein engineering studies and may give an indication of what the protein can tolerate in terms of mutagenesis without losing function.

Manipulation of the HA nucleic acid molecules of the invention may result in the production of HA proteins having functional differences from the wild-type HA proteins. These proteins may be improved in efficiency or activity, may be present in greater numbers in the cell than is usual, or may be decreased in efficiency or activity.

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The modulation of activity or number of HA proteins involved in cell wall biosynthesis or rearrangements may both impact the production, yield, and/or efficiency of production of one or more fine chemicals from C glutamicum cells. For example, by altering the activity of these proteins, it may be possible to modulate the structure or thickness of the cell wall. The cell wall serves in large measure as a protective device against osmotic lysis and external sources of injury; by modifying the cell wall it may be possible to increase the ability of C. glutamicum to withstand the mechanical and shear force stresses encountered by this microorganism during large-scale fermentor culture. Further, each C. glutamicum cell is surrounded by a thick cell wall, and thus, a significant portion of the biomass present in large scale culture consists of cell wall. By increasing the rate at which the cell wall is synthesized or by activating cell wall synthesis (through genetic engineering of the HA cell wall proteins of the invention) it may be possible to improve the growth rate of the microorganism. Similarly, by decreasing the activity or number of proteins involved in the degradation of cell wall or by decreasing the repression of cell wall biosynthesis, an overall increase in cell wall production may be achieved. An increase in the number of viable C. glutumicum cells (as may be accomplished by any of the foregoing described protein alterations) should result in increased numbers of cells producing the desired fine chemical in large-scale fermentor culture, which should permit increased yields or efficiency of production of these compounds from the culture.

The modulation of activity or number of C. glutamicum HA proteins that participate in the modification or degradation of aromatic or aliphatic compounds may also have direct or indirect impacts on the production of one or more fine chemicals from these cells. Certain aromatic or aliphatic modification or degradation products are desirable fine chemicals (e.g., organic acids or modified aromatic and aliphatic compounds); thus, by modifying the enzymes which perform these modifications (e.g., hydroxylation, methylation, or isomerization) or degradation reactions, it may be possible to increase the yields of these desired compounds. Similarly, by decreasing the activity or number of proteins involved in pathways which further degrade the modified

or breakdown products of the aforementioned reactions it may be possible to improve the yields of these fine chemicals from C glutamicum cells in culture.

These aromatic and aliphatic modification and degradative enzymes are themselves fine chemicals. In purified form, these enzymes may be used to degrade aromatic and aliphatic compounds (e.g., toxic chemicals such as petroleum products), either for the bioremediation of polluted sites, for the engineered decomposition of wastes, or for the large-scale and economically feasible production of desired modified aromatic or aliphatic compounds or their breakdown products, some of which may be conveniently used as carbon or energy sources for other fine chemical-producing compounds in culture (see, e.g., Faber, K. (1995) Biotransformations in Organic Chemistry, Springer: Berlin and references therein; and Roberts, S.M., ed. (1992-1996) Preparative Biotransformations, Wiley: Chichester, and references therein). By genetically altering these proteins such that their regulation by other cellular mechanisms is lessened or abolished, it may be possible to increase the overall number or activity of these proteins, thereby improving not only the yield of these fine chemicals but also the activity of these harvested proteins.

The modification of these aromatic and aliphatic modifying and degradation enzymes may also have an indirect effect on the production of one or more fine chemical. Many aromatic and aliphatic compounds (such as those that may be encountered as impurities in culture media or as waste products from cellular metabolism) are toxic to cells; by modifying and/or degrading these compounds such that they may be readily removed or destroyed, cellular viability should be increased. Further, these enzymes may modify or degrade these compounds in such a manner that the resulting products may enter the normal carbon metabolism pathways of the cell, thus rendering the cell able to use these compounds as alternate carbon or energy sources. In large-scale culture situations, when there may be limiting amounts of optimal carbon sources, these enzymes provide a method by which cells may continue to grow and divide using aromatic or aliphatic compounds as nutrients. In either case, the resulting increase in the number of *C. glutumicum* cells in the culture producing the desired fine chemical should in turn result in increased yields or efficiency of production of the fine chemical(s).

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Modifications in activity or number of HA proteins involved in the metabolism of inorganic compounds may also directly or indirectly affect the production of one or more fine chemicals from C. glutamicum or related bacterial cultures. For example, many desirable fine chemicals, such as nucleic acids, amino acids, cofactors and vitamins (e.g., thiamine, biotin, and lipoic acid) cannot be synthesized without inorganic molecules such as phosphorous, nitrate, sulfate, and iron. The inorganic metabolism

proteins of the invention permit the cell to obtain these molecules from a variety of inorganic compounds and to divert them into various fine chemical biosynthetic pathways. Therefore, by increasing the activity or number of enzymes involved in the metabolism of these inorganic compounds, it may be possible to increase the supply of these possibly limiting inorganic molecules, thereby directly increasing the production or efficiency of production of various fine chemicals from C. glutanicum cells containing such altered proteins. Modification of the activity or number of inorganic metabolism enzymes of the invention may also render C. glutanicum able to better utilize limited inorganic compound supplies, or to utilize nonoptimal inorganic compounds to synthesize amino acids, vitamins, cofactors, or nucleic acids, all of which are necessary for continued growth and replication of the cell. By improving the viability of these cells in large-scale culture, the number of C. glutonicum cells producing one or more fine chemicals in the culture may also be increased, in turn increasing the yields or efficiency of production of one or more fine chemicals.

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C. glutamicum enzymes for general processes are themselves desirable fine chemicals. The specific properties of enzymes (i.e., regio- and stereospecificity, among others) make them useful catalysts for chemical reactions in vitro. Either whole C. glutamicum cells may be incubated with an appropriate substrate such that the desired product is produced by enzymes in the cell, or the desired enzymes may be overproduced and purified from C. glutamicum cultures (or those of a related bacterium) and subsequently utilized in in vitro reactions in an industrial setting (either in solution or immobilized on a suitable immobile phase). In either situation, the enzyme can either be a natural C glutamicum protein, or it may be mutagenized to have an altered activity; typical industrial uses for such enzymes include as catalysts in the chemical industry (e.g., for synthetic organic chemistry) as food additives, as feed components, for fruit processing, for leather preparation, in detergents, in analysis and medicine, and in the textile industry (see, e.g., Yamada, H. (1993) "Microbial reactions for the production of useful organic compounds," Chimica 47: 5-10; Roberts, S.M. (1998) Preparative biotransformations: the employment of enzymes and whole-cells in synthetic chemistry," J. Chem. Soc. Perkin Trans. 1: 157-169; Zaks, A. and Dodds, D.R. (1997) "Application of biocatalysis and biotransformations to the synthesis of pharmaceuticals," DDT 2: 513-531; Roberts, S.M. and Williamson, N.M. (1997) "The use of enzymes for the preparation of biologically active natural products and analogues in optically active form," Curr. Organ Chemistry 1: 1-20; Faber, K. (1995) Biotransformations in Organic Chemistry, Springer: Berlin; Roberts, S.M., ed. (1992-96) Preparative Biotransformations, Wiley: Chichester; Cheetham, P.S.J. (1995) "The applications of enzymes in industry" in : Handbook of Enzyme Biotechnology, 3rd ed.,

Wiseman, A., ed., Elis: Horwood, p. 419-552; and Ullmann's Encyclopedia of Industrial Chemistry (1987), vol. A9, Enzymes, p. 390-457). Thus, by increasing the activity or number of these enzymes, it may be possible to also increase the ability of the cell to convert supplied substrates to desired products, or to overproduce these enzymes for increased yields in large-scale culture. Further, by mutagenizing these proteins it may be possible to remove feedback inhibition or other repressive cellular regulatory controls such that greater numbers of these enzymes may be produced and activated by the cell, thereby leading to greater yields, production, or efficiency of production of these fine chemical proteins from large-scale cultures. Further, manipulation of these enzymes may alter the activity of one or more C. glutamicum metabolic pathways, such as those for the biosynthesis or secretion of one or more fine chemicals.

Mutagenesis of the proteolytic enzymes of the invention such that they are altered in activity or number may also directly or indirectly affect the yield, production, and/or efficiency of production of one or more fine chemicals from C. glutamicum. For example, by increasing the activity or number of these proteins, it may be possible to increase the ability of the bacterium to survive in large-scale culture, due to an increased ability of the cell to rapidly degrade proteins misfolded in response to the high temperatures, nonoptimal pH, and other stresses encountered during fermentor culture. Increased numbers of cells in these cultures may result in increased yields or efficiency of production of one or more desired fine chemicals, due to the relatively larger number of cells producing these compounds in the culture. Also, C glutamicum cells possess multiple cell-surface proteases which serve to break down external nutrients into molecules which may be more readily incorporated by the cells as carbon/energy sources or nutrients of other kinds. An increase in activity or number of these enzymes may improve this turnover and increase the levels of available nutrients, thereby improving cell growth or production. Thus, modifications of the proteases of the invention may indirectly impact C. glutamicum fine chemical production.

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A more direct impact on fine chemical production in response to the modification of one or more of the proteases of the invention may occur when these proteases are involved in the production or degradation of a desired fine chemical. By decreasing the activity of a protease which degrades a fine chemical or a protein involved in the synthesis of a fine chemical it may be possible to increase the levels of that fine chemical (due to the decreased degradation or increased synthesis of the compound). Similarly, by increasing the activity of a protease which degrades a compound to result in a fine chemical or a protein involved in the degradation of a fine chemical, a similar result should be achieved: increased levels of the desired fine chemical from *C. glutamicum* cells containing these engineered proteins.

The aforementioned mutagenesis strategies for HA proteins to result in increased yields of a fine chemical from C glutamicum are not meant to be limiting; variations on these strategies will be readily apparent to one skilled in the art. Using such strategies, and incorporating the mechanisms disclosed herein, the nucleic acid and protein molecules of the invention may be utilized to generate C glutamicum or related strains of bacteria expressing mutated HA nucleic acid and protein molecules such that the yield, production, and/or efficiency of production of a desired compound is improved. This desired compound may be any product produced by C. glutamicum, which includes the final products of biosynthesis pathways and intermediates of naturally-occurring metabolic pathways, as well as molecules which do not naturally occur in the metabolism of C glutamicum, but which are produced by a C. glutamicum strain of the invention.

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This invention is further illustrated by the following examples which should not be construed as limiting. The contents of all references, patent applications, patents, and published patent applications cited throughout this application are hereby incorporated by reference.

### Exemplification

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### Example 1: Preparation of total genomic DNA of Corynebacterium glutamicum ATCC 13032

A culture of Corynebacterium glutamicum (ATCC 13032) was grown overnight at 30°C with vigorous shaking in BHI medium (Difco). The cells were harvested by centrifugation, the supernatant was discarded and the cells were resuspended in 5 ml buffer-I (5% of the original volume of the culture — all indicated volumes have been calculated for 100 ml of culture volume). Composition of buffer-1: 140.34 g/l sucrose, 2.46 g/l MgSO, x 7H2O, 10 ml/l KH2PO, solution (100 g/l, adjusted to pH 6.7 with KOH), 50 ml/l M12 concentrate (10 g/l (NH.)2SO., 1 g/l NaCl, 2 g/l MgSO, x 7H2O, 0.2 g/l CaCl<sub>2</sub>, 0.5 g/l yeast extract (Difco), 10 ml/l trace-elements-mix (200 mg/l FeSO, x H<sub>2</sub>O, 10 mg/l ZnSO, x 7 H<sub>2</sub>O, 3 mg/l MnCl<sub>2</sub> x 4 H<sub>2</sub>O, 30 mg/l H<sub>3</sub>BO, 20 mg/l CoCl<sub>2</sub> x 6 H<sub>2</sub>O, 1 mg/l NiCl<sub>2</sub> x 6 H<sub>2</sub>O, 3 mg/l Na<sub>2</sub>MoO, x 2 H<sub>2</sub>O, 500 mg/l complexing agent (EDTA or critic acid), 100 ml/l vitamins-mix (0.2 mg/l biotin, 0.2 mg/l folic acid, 20 mg/l p-amino benzoic acid, 20 mg/l riboflavin, 40 mg/l ca-panthothenate, 140 mg/l nicotinic acid, 40 mg/l pyridoxole hydrochloride, 200 mg/l myo-inositol). Lysozyme was added to the suspension to a final concentration of 2.5 mg/ml. After an approximately 4 h incubation at 37°C, the cell wall was degraded and the resulting protoplasts are harvested by centrifugation. The pellet was washed once with 5 ml buffer-I and once with 5 ml TE-buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8). The pellet was resuspended in 4 ml TE-buffer and 0.5 ml SDS solution (10%) and 0.5 ml NaCl solution (5 M) are added. After adding of proteinase K to a final concentration of 200  $\mu$ g/ml, the suspension is incubated for ca.18 h at 37°C. The DNA was purified by extraction with phenol, phenol-chloroform-isoamylalcohol and chloroformisoamylalcohol using standard procedures. Then, the DNA was precipitated by adding 1/50 volume of 3 M sodium acetate and 2 volumes of ethanol, followed by a 30 min incubation at -20°C and a 30 min centrifugation at 12,000 rpm in a high speed centrifuge using a SS34 rotor (Sorvall). The DNA was dissolved in 1 ml TE-buffer containing 20

μg/ml RNaseA and dialysed at 4°C against 1000 ml TE-buffer for at least 3 hours. During this time, the buffer was exchanged 3 times. To aliquots of 0.4 ml of the dialysed DNA solution, 0.4 ml of 2 M LiCl and 0.8 ml of ethanol are added. After a 30 min incubation at -20°C, the DNA was collected by centrifugation (13,000 rpm, Biofuge Fresco, Heraeus, Hanau, Germany). The DNA pellet was dissolved in TE-buffer. DNA prepared by this procedure could be used for all purposes, including southern blotting or construction of genomic libraries.

### Example 2: Construction of genomic libraries in Escherichia coli of Corynebacterium glutamicum ATCC13032.

Starting from DNA prepared as described in Example 1, cosmid and plasmid libraries were constructed according to known and well established methods (see e.g., Sambrook, J. et al. (1989) "Molecular Cloning: A Laboratory Manual", Cold Spring Harbor Laboratory Press, or Ausubel, F.M. et al. (1994) "Current Protocols in Molecular Biology", John Wiley & Sons.)

Any plasmid or cosmid could be used. Of particular use were the plasmids pBR322 (Sutcliffe, J.G. (1979) Proc. Natl. Acad. Sci. USA, 75·3737-3741), pACYC177 (Change & Cohen (1978) J. Bacteriol 134:1141-1156), plasmids of the pBS series (pBSSK+, pBSSK- and others; Stratagene, LaJolla, USA), or cosmids as SuperCos1 (Stratagene, LaJolla, USA) or Loristó (Gibson, T.J., Rosenthal A. and Waterson, R.H. (1987) Gene 53:283-286.

### Example 3: DNA Sequencing and Computational Functional Analysis

Genomic libraries as described in Example 2 were used for DNA sequencing according to standard methods, in particular by the chain termination method using ABI377 sequencing machines (see e.g., Fleischman, R.D. et al. (1995) "Whole-genome Random Sequencing and Assembly of Haemophilus Influenzae Rd., Science, 269:496-512). Sequencing primers with the following nucleotide sequences were used: 5'-GGAAACAGTATGACCATG-3' or 5'-GTAAAACGACGGCCAGT-3'.

### 30 Example 4: In vivo Mutagenesis

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In vivo mutagenesis of Corynebacterium glutamicum can be performed by passage of plasmid (or other vector) DNA through E coli or other microorganisms (e.g. Bacillus spp. or yeasts such as Sacuharomyces cerevisiae) which are impaired in their capabilities to maintain

the integrity of their genetic information. Typical mutator strains have mutations in the genes for the DNA repair system (e.g., mutHLS, mutD, mutT, etc.; for reference, see Rupp, W.D. (1996) DNA repair mechanisms, in: *Escherichia col*i and *Salmonella*, p. 2277-2294, ASM: Washington.) Such strains are well known to those skilled in the art. The use of such strains is illustrated, for example, in Greener, A. and Callahan, M. (1994) Strategies 7: 32-34.

### Example 5: DNA Transfer Between Escherichia coli and Corynebacterium glutamicum

Several Corynebacterium and Brevibacterium species contain endogenous plasmids (as e.g., pHM1519 or pBL1) which replicate autonomously (for review see, e.g., Martin, J.F. et al. (1987) Biotechnology, 5:137-146). Shuttle vectors for Escherichia coli and Corynebacterium glutamicum can be readily constructed by using standard vectors for E coli (Sambrook, J. et al. (1989), "Molecular Cloning: A Laboratory Manual", Cold Spring Harbor Laboratory Press or Ausubel, F.M. et al (1994) "Current Protocols in Molecular Biology", John Wiley & Sons) to which a origin or replication for and a suitable marker from Corynebacterium glutumicum is added. Such origins of replication are preferably taken from endogenous plasmids isolated from Corynebacterium and Brevibacterium species. Of particular use as transformation markers for these species are genes for kanamycin resistance (such as those derived from the Tn5 or Tn903 transposons) or chloramphenicol (Winnacker, E.L. (1987) "From Genes to Clones -Introduction to Gene Technology, VCH, Weinheim). There are numerous examples in the literature of the construction of a wide variety of shuttle vectors which replicate in both E. coli and C. glutamicum, and which can be used for several purposes, including gene overexpression (for reference, see e.g., Yoshihama, M. et al. (1985) J. Bacteriol. 162:591-597, Martin J.F. et al. (1987) Biotechnology, 5:137-146 and Eikmanns, B.J. et al. (1991) Gene, 102:93-98).

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Using standard methods, it is possible to clone a gene of interest into one of the shuttle vectors described above and to introduce such a hybrid vectors into strains of Corynebacterium glutamicum. Transformation of C. glutamicum can be achieved by protoplast transformation (Kastsumata, R. et al. (1984) J. Bacteriol. 159306-311), electroporation (Liebl, E. et al. (1989) FEMS Microbiol. Letters, 53:399-303) and in cases where special vectors are used, also by conjugation (as described e.g. in Schäfer, A et al. (1990) J. Bacteriol. 172:1663-1666). It is also possible to transfer the shuttle vectors for



C glutamicum to E. coli by preparing plasmid DNA from C' glutamicum (using standard methods well-known in the art) and transforming it into E. coli. This transformation step can be performed using standard methods, but it is advantageous to use an Mcr-deficient E. coli strain, such as NM522 (Gough & Murray (1983) J. Mol. Biol. 166:1-19).

### Example 6: Assessment of the Expression of the Mutant Protein

Observations of the activity of a mutated protein in a transformed host cell rely on the fact that the mutant protein is expressed in a similar fashion and in a similar quantity to that of the wild-type protein. A useful method to ascertain the level of transcription of the mutant gene (an indicator of the amount of mRNA available for translation to the gene product) is to perform a Northern blot (for reference see, for example, Ausübel et al. (1988) Current Protocols in Molecular Biology, Wiley: New York), in which a primer designed to bind to the gene of interest is labeled with a detectable tag (usually radioactive or chemiluminescent), such that when the total RNA of a culture of the organism is extracted, run on gel, transferred to a stable matrix and incubated with this probe, the binding and quantity of binding of the probe indicates the presence and also the quantity of mRNA for this gene. This information is evidence of the degree of transcription of the mutant gene. Total cellular RNA can be prepared from Corynebacterium glutamicum by several methods, all well-known in the art, such as that described in Bormann, E.R. et al. (1992) Mol. Microbiol. 6: 317-326.

To assess the presence or relative quantity of protein translated from this mRNA, standard techniques, such as a Western blot, may be employed (see, for example, Ausubel et al. (1988) Current Protocols in Molecular Biology, Wiley: New York). In this process, total cellular proteins are extracted, separated by gel electrophoresis, transferred to a matrix such as nitrocellulose, and incubated with a probe, such as an antibody, which specifically binds to the desired protein. This probe is generally tagged with a chemiluminescent or colorimetric label which may be readily detected. The presence and quantity of label observed indicates the presence and quantity of the desired mutant protein present in the cell.

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### Example 7: Growth of Genetically Modified Corynebacterium glutamicum — Media and Culture\_Conditions

Genetically modified Corynebacteria are cultured in synthetic or natural growth media. A number of different growth media for Corynebacteria are both well-known and readily available (Lieb et al. (1989) Appl. Microbiol. Biotechnol., 32:205-210; von der Osten et al. (1998) Biotechnology Letters, 11:11-16; Patent DE 4,120,867; Liebl (1992) "The Genus Corynebacterium, in: The Procaryotes, Volume II, Balows, A. et al., eds. Springer-Verlag). These media consist of one or more carbon sources, nitrogen sources, inorganic salts, vitamins and trace elements. Preferred carbon sources are sugars, such as mono-, di-, or polysaccharides. For example, glucose, fructose, mannose, galactose, ribose, sorbose, ribulose, lactose, maltose, sucrose, raffinose, starch or cellulose serve as very good carbon sources. It is also possible to supply sugar to the media via complex compounds such as molasses or other by-products from sugar refinement. It can also be advantageous to supply mixtures of different carbon sources. Other possible carbon sources are alcohols and organic acids, such as methanol, ethanol, acetic acid or lactic 15 acid. Nitrogen sources are usually organic or inorganic nitrogen compounds, or materials which contain these compounds. Exemplary nitrogen sources include ammonia gas or ammonia salts, such as NH,Cl or (NH,),SO,, NH,OH, nitrates, urea, amino acids or complex nitrogen sources like com steep liquor, soy bean flour, soy bean protein, yeast extract, meat extract and others. 20

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Inorganic salt compounds which may be included in the media include the chloride-, phosphorous- or sulfate- salts of calcium, magnesium, sodium, cobalt, molybdenum, potassium, manganese, zinc, copper and iron. Chelating compounds can be added to the medium to keep the metal ions in solution. Particularly useful chelating compounds include dihydroxyphenols, like catechol or protocatechuate, or organic acids, such as citric acid. It is typical for the media to also contain other growth factors, such as vitamins or growth promoters, examples of which include biotin, riboflavin, thiamin, folic acid, nicotinic acid, pantothenate and pyridoxin. Growth factors and salts frequently originate from complex media components such as yeast extract, molasses, corn steep liquor and others. The exact composition of the media compounds depends strongly on the immediate experiment and is individually decided for each specific case. Information about media optimization is available in the textbook "Applied Microbiol. Physiology, A Practical Approach (eds. P.M. Rhodes, P.F. Stanbury, IRL Press (1997) pp. 53-73, ISBN 0

19 963577 3). It is also possible to select growth media from commercial suppliers, like standard 1 (Merck) or BHI (grain heart infusion, DIFC) or others.

All medium components are sterilized, either by heat (20 minutes at 1.5 bar and 121°C) or by sterile filtration. The components can either be sterilized together or, if necessary, separately. All media components can be present at the beginning of growth, or they can optionally be added continuously or batchwise.

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Culture conditions are defined separately for each experiment. The temperature should be in a range between 15°C and 45°C. The temperature can be kept constant or can be altered during the experiment. The pH of the medium should be in the range of 5 to 8.5, preferably around 7.0, and can be maintained by the addition of buffers to the media. An exemplary buffer for this purpose is a potassium phosphate buffer. Synthetic buffers such as MOPS, HEPES, ACES and others can alternatively or simultaneously be used. It is also possible to maintain a constant culture pH through the addition of NaOH or NH,OH during growth. If complex medium components such as yeast extract are utilized, the necessity for additional buffers may be reduced, due to the fact that many complex compounds have high buffer capacities. If a fermentor is utilized for culturing the microorganisms, the pH can also be controlled using gaseous ammonia.

The incubation time is usually in a range from several hours to several days. This time is selected in order to permit the maximal amount of product to accumulate in the broth. The disclosed growth experiments can be carried out in a variety of vessels, such as microtiter plates, glass tubes, glass flasks or glass or metal fermentors of different sizes. For screening a large number of clones, the microorganisms should be cultured in microtiter plates, glass tubes or shake flasks, either with or without baffles. Preferably 100 ml shake flasks are used, filled with 10% (by volume) of the required growth medium. The flasks should be shaken on a rotary shaker (amplitude 25 mm) using a speed-range of 100 – 300 rpm. Evaporation losses can be diminished by the maintenance of a humid atmosphere; alternatively, a mathematical correction for evaporation losses should be performed.

If genetically modified clones are tested, an unmodified control clone or a control clone containing the basic plasmid without any insert should also be tested. The medium is inoculated to an OD<sub>600</sub> of O.5 – 1.5 using cells grown on agar plates, such as CM plates (10 g/l glucose, 2,5 g/l NaCl, 2 g/l urea, 10 g/l polypeptone, 5 g/l yeast extract, 5 g/l meat extract, 22 g/l NaCl, 2 g/l urea, 10 g/l polypeptone, 5 g/l yeast extract, 5 g/l meat extract,

22 g/l agar, pH 6.8 with 2M NaOH) that had been incubated at 30°C. Inoculation of the media is accomplished by either introduction of a saline suspension of C glutamicum cells from CM plates or addition of a liquid preculture of this bacterium.

### 5 Example 8 - In vitro Analysis of the Function of Mutant Proteins

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The determination of activities and kinetic parameters of enzymes is well established in the art. Experiments to determine the activity of any given altered enzyme must be tailored to the specific activity of the wild-type enzyme, which is well within the ability of one skilled in the art. Overviews about enzymes in general, as well as specific details concerning structure, kinetics, principles, methods, applications and examples for the determination of many enzyme activities may be found, for example, in the following references: Dixon, M., and Webb, E.C., (1979) Enzymes. Longmans: London; Fersht, (1985) Enzyme Structure and Mechanism. Freeman: New York; Walsh, (1979) Enzymatic Reaction Mechanisms. Freeman: San Francisco; Price, N.C., Stevens, L. (1982) Fundamentals of Enzymology. Oxford Univ. Press: Oxford; Boyer, P.D., ed. (1983) The Enzymes, 3<sup>rd</sup> ed. Academic Press: New York; Bisswanger, H., (1994) Enzymkinetik, 2<sup>rd</sup> ed. VCH: Weinheim (ISBN 3527300325); Bergmeyer, H.U., Bergmeyer, J., Graßl, M., eds. (1983-1986) Methods of Enzymatic Analysis, 3<sup>rd</sup> ed., vol. 1-XII, Verlag Chemie: Weinheim; and Ullmann's Encyclopedia of Industrial Chemistry (1987) vol. A9, "Enzymes". VCH: Weinheim, p. 352-363.

The activity of proteins which bind to DNA can be measured by several well-established methods, such as DNA band-shift assays (also called gel retardation assays). The effect of such proteins on the expression of other molecules can be measured using reporter gene assays (such as that described in Kolmar, H. et al. (1995) EMBO J. 14: 3895-3904 and references cited therein). Reporter gene test systems are well known and established for applications in both pro- and eukaryotic cells, using enzymes such as beta-galactosidase, green fluorescent protein, and several others.

The determination of activity of membrane-transport proteins can be performed according to techniques such as those described in Gennis, R.B. (1989) "Pores, Channels and Transporters", in Biomembranes, Molecular Structure and Function, Springer: Heidelberg, p. 85-137; 199-234; and 270-322.

### Example 9: Analysis of Impact of Mutant Protein on the Production of the Desired Product

The effect of the genetic modification in C. glutamicum on production of a desired compound (such as an amino acid) can be assessed by growing the modified microorganism under suitable conditions (such as those described above) and analyzing

the medium and/or the cellular component for increased production of the desired product (i.e., an amino acid). Such analysis techniques are well known to one skilled in the art, and include spectroscopy, thin layer chromatography, staining methods of various kinds, enzymatic and microbiological methods, and analytical chromatography such as high performance liquid chromatography (see, for example, Ullman, Encyclopedia of Industrial Chemistry, vol. A2, p. 89-90 and p. 443-613, VCH: Weinheim (1985); Fallon, A. et al., (1987) "Applications of HPLC in Biochemistry" in: Laboratory Techniques in Biochemistry and Molecular Biology, vol. 17; Rehm et al. (1993) Biotechnology, vol. 3, Chapter III: "Product recovery and purification", page 469-714, VCH: Weinheim; Belter, P.A. et al. (1988) Bioseparations: downstream processing for biotechnology, John Wiley and Sons; Kennedy, J.F. and Cabral, J.M.S. (1992) Recovery processes for biological materials, John Wiley and Sons; Shaeiwitz, J.A. and Henry, J.D. (1988) Biochemical separations, in: Ulmann's Encyclopedia of Industrial Chemistry, vol. B3, Chapter 11, page 1-27, VCH: Weinheim; and Dechow, F.J. (1989) Separation and purification techniques in biotechnology, Noyes Publications.)

In addition to the measurement of the final product of fermentation, it is also possible to analyze other components of the metabolic pathways utilized for the production of the desired compound, such as intermediates and side-products, to determine the overall efficiency of production of the compound. Analysis methods include measurements of nutrient levels in the medium (e.g., sugars, hydrocarbons, nitrogen sources, phosphate, and other ions), measurements of biomass composition and growth, analysis of the production of common metabolites of biosynthetic pathways, and measurement of gasses produced during fermentation. Standard methods for these measurements are outlined in Applied Microbial Physiology, A Practical Approach, P.M. Rhodes and P.F. Stanbury, eds., IRL Press, p. 103-129; 131-163; and 165-192 (ISBN: 0199635773) and references cited therein.

### Example 10: Purification of the Desired Product from C. glutamicum Culture

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Recovery of the desired product from the *C glutamicum* cells or supernatant of the above-described culture can be performed by various methods well known in the art. If the desired product is not secreted from the cells, the cells can be harvested from the culture by low-speed centrifugation, the cells can be lysed by standard techniques, such as mechanical force or sonication. The cellular debris is removed by centrifugation, and the supernatant fraction containing the soluble proteins is retained for further purification of the desired compound. If the product is secreted from the *C. glutamicum* 

cells, then the cells are removed from the culture by low-speed centrifugation, and the supernate fraction is retained for further purification.

The supernatant fraction from either purification method is subjected to chromatography with a suitable resin, in which the desired molecule is either retained on a chromatography resin while many of the impurities in the sample are not, or where the impurities are retained by the resin while the sample is not. Such chromatography steps may be repeated as necessary, using the same or different chromatography resins. One skilled in the art would be well-versed in the selection of appropriate chromatography resins and in their most efficacious application for a particular molecule to be purified. The purified product may be concentrated by filtration or ultrafiltration, and stored at a temperature at which the stability of the product is maximized.

There are a wide array of purification methods known to the art and the preceding method of purification is not meant to be limiting. Such purification techniques are described, for example, in Bailey, J.E. & Ollis, D.F. Biochemical Engineering Fundamentals, McGraw-Hill: New York (1986).

The identity and purity of the isolated compounds may be assessed by techniques standard in the art. These include high-performance liquid chromatography (HPLC), spectroscopic methods, staining methods, thin layer chromatography, NIRS, enzymatic assay, or microbiologically. Such analysis methods are reviewed in: Patek et al. (1994) Appl Environ Microbiol. 60: 133-140; Malakhova et al. (1996) Biotekhnologiya 11: 27-32; and Schmidt et al. (1998) Bioprocess Engineer. 19: 67-70. Ulmann's Encyclopedia of Industrial Chemistry, (1996) vol. A27, VCH: Weinheim, p. 89-90, p. 521-540, p. 540-547, p. 559-566, 575-581 and p. 581-587, Michal, G. (1999) Biochemical Pathways: An Atlas of Biochemistry and Molecular Biology, John Wiley and Sons; Fallon, A. et al. (1987) Applications of HPLC in Biochemistry in: Laboratory Techniques in Biochemistry and Molecular Biology, vol. 17.

### Equivalents

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Those skilled in the art will recognize, or will be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims.

## Table 1: Genes in the Application

### Cell wall biosynthesis

			•		
Function	N. ACETYLMURAMOYL-L. ALANINE AMIDASE (EC 15.128) N. ACETYLMURAMOYL-L. ALANINE AMIDASE (EC 35.128) N. ACETYLLMURAMOYL-L. ALANINE AMIDASE (EC 35.128) UDP.N. ACETYLGLUCOSAMINE I CARBOXYVINYLTRANSFERASE (EC 2.51.1) PHOSPHO.N. ACETYLMURAMATE.—ALANINE LIGASE (EC 6.12.8) UDP.N. ACETYLMURAMATE.—ALANINE LIGASE (EC 6.12.9) UDP.N. ACETYLMURAMOYLALANINE. D. GLUTAMATE LIGASE (EC 6.12.9) UDP.N. ACETYLMURAMOYLALANINE. D. GLUTAMATE. 2,6 DIAMINOPIMELATE LIGASE (EC 6.12.13) UDP.N. ACETYLMURAMOYLALANYL. D. GLUTAMAYL. 2,6 DIAMINOPIMELATE. D. ALANYL. D. ALANYL LIGASE (EC 6.12.15)	UDP-N-ACETYLMURAMOYLALAN IL-D-OLU TAMYL-2,6-DIAMINOPIMELATE-D-ALANYL-D-ALANYL LIGASE UDP-N-ACETYLMURAMOYLALAN IL-D-GLUTAMYL-2,6-DIAMINOPIMELATE-D-ALANYL-D-ALANYL LIGASE (EC 8.3.2.15)	UDP-N. ACETYLMURAMOYLALANYL, O-GLUTAMATE, 2,6-DIAMINOPIMELATE LICASE (EC. 5, 2, 1, 2) ALANINE RACEMASE (EC. 5, 1, 1, 1) D. ALANINED-ALANINE LIGASE (EC. 8, 3, 2, 4) UDP-N. ACETYLGLUCOSAMINEN-ACETYLGLUCOSAMINE TRANSFERASE (EC. 2, 4, 1, 4) UNDECAPRENOL N. ACETYLGLUCOSAMINE TRANSFERASE (EC. 2, 4, 1, 4)	PENICILLIN-BINDING PROTEIN 2 PENICILLIN-BINDING PROTEIN 5 PENICILLIN-BINDING PROTEIN 4 PENICILLIN-BINDING PROTEIN 14 PENICILLIN-BINDING PROTEIN 1A PENICILLIN-BINDING PROTEIN 1A PENICILLIN-BINDING PROTEIN 1A PENICILLIN-BINDING PROTEIN 3 PENICILLIN-BINDING PROTEIN 1A PEC 3 4, 19 4, 1 D. ALANINE-ENDOPEPTIDASE (EC 3 4, 99 -)	(ALCO9883) penecilin binding protein (Mycobacterium troerculosis) Glycosyllransferases, typically involved in cell wall biogenesis perosamine synthelase
<b>Gene Name</b>	BS-murA,EC-murA EC-mraY,BS-mraY EC-mwC,BS-murC BS-radE,EC-murl BS-mwD	BS-murF	BS-murE, EC-murE BS-yncD, EC-alr EC-ddA, BS-ddIA EC-murG, BS-murG	EC-fist, BS-spoVD BS-pbpF, EC-mrcA BS-pbpC	BS-yvfE,EC-b2253
NT Stop	8271 3022 2962 5813 115 997 4388 2539 8920	7260	8473 1921 806 1610	10162 121 4853 4457 6315 1187 16650	2675 3759 20498
Start	7458 5097 1709 6910 1572 1845 5803 7264	7694 8451	10035 1193 3 2698	12273 846 3928 3525 7716 3	837 2872 21652
Config	GR00417 GR00749 GR00758 GR00758 GR00758 GR00758	GR00758 GR00758	GR00758 GR00127 GR00292 GR00758	GR00758 GR0162 GR0152 GR0158 GR00158 GR00162 GR00162	GR00449 GR00400 GR00367
Identification Code	RXA01430 RXA02641 RXA02135 RXA02708 RXA02702 RXA02411 RXA02705 RXA01254 RXA01254	RXA02708	RXA02710 RXA00508 RXA01022 RXA02703	RXA02711 RXA02859 RXA00569 RXA00594 RXA01828 RXA01612 RXA01612	RXA01608 RXA01376 RXA01270

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Function	CELL DIVISION PROTEIN FTSW CELL DIVISION PROTEIN FTSZ CELL DIVISION PROTEIN FTSX CELL DIVISION ATP-BINDING PROTEIN FTSE CELL DIVISION ATP-BINDING PROTEIN FTSE CELL DIVISION ROTEIN FTSK CELL DIVISION PROTEIN FTSK CELL DIVISION PROTEIN FTSK CELL DIVISION CONTROL PROTEIN 15 (EC 2 7 1 -) CELL CYCLE PROTEIN MESJ CELL DIVISION PROTEIN FTSK CELL DIVISION PROTEIN FTSK CELL DIVISION PROTEIN FTSK CELL DIVISION PROTEIN FTSK CELL DIVISION PROTEIN FTSY Hypothelical Cell Division Prolein miaW	F1SU RSH suppressor - suppressors the temperature-sensitivity of the fish I(Ts) mutation GLUCOSE INHIBITED DIVISION PROTEIN B CHROMOSOME SEGREGATION PROTEIN SMC2 STAGE 0 SPORULATION PROTEIN J STAGE II SPORULATION PROTEIN E STAGE V SPORULATION PROTEIN E SOJ PROTEIN	Function	METHIONINE AMINOPEPTIDASE (EC 3 4 11 18) METHIONINE AMINOPEPTIDASE (EC 3 4 11 18) ATP-DEPENDENT PROTEASE LA (EC 3 4 21 53) ATP-DEPENDENT PROTEASE LA (EC 3 4 21 53) ATP-DEPENDENT PROTEASE LA (EC 3 4 21 53) ATP-DEPENDENT PROTEASE LA (EC 3 4 24 -) ZINC METALLOPROTEASE (EC 3 4 24 -) ZINC METALLOPROTEASE (EC 3 4 24 -) ZINC METALLOPROTEASE (EC 3 4 24 -) ATP-DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPA ATP-DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT
Gene Name	BS-spove.EC-flsW BS-flsZ.EC-flsZ BS-flsE,EC-flsE EC-b2304,BS-yMF BS-spollIE,EC-flsK BS-flsH,EC-hIIB BS-flsH,EC-hIIB BS-flsY BS-yxA,EC-yabC	EC-sMB.BS-ylyB BS-gidB,EC-gidB BS-smc BS-sp00J BS-yukA	Gene Nаme	EC-map, BS-map EC-yifB BS-yibL BS-yibC BS-hirA EC-hirA
Slop	2694 646 646 1562 1862 5 5 1291 1751 871 17596 20926 5906	2984 2970 3403 17154 5631 1344 7736 4432 14683	Stop	484 3612 8857 2176 1981 30 1052 3196 4991 1332 497
Slart	4382 2729 1545 2248 6328 6328 1588 2 2 2 2 16655 18388 4161	3480 2041 2777 14248 4495 4661 9058 3512	Slart	2 2740 5337 3225 386 1640 1954 2216 3159 2654
Contig	GR00758 GR00759 GR00002 GR00002 GR00023 GR00234 GR00424 GR00424 GR00424	GR00759 GR00752 GR00717 GR00417 GR00417 GR00417 GR00417 GR00417	Contig	GR00178 GR00449 GR00593 GR00459 GR00534 GR00514 GR00715 GR00715
Identification	RXA02704 RXA02722 RXA00009 RXA000143 RXA000143 RXA00857 RXA00857 RXA01435 RXA01511 RXA01513	RXA02713 RXA02651 RXA02651 RXA01426 RXA01428 RXA01640 RXA01640 RXA01603	Proteolysis Identification Code	RXA00675 RXA01609 RXA01358 RXA01458 RXA01654 RXA01669 RXA01669 RXA01669 RXA02470 RXA02630
		•		

			XYPEPTIDASE) (EC 34 164)				
Function	PROBABLE PERIPLASMIC SERINE PROTEASE UC-LINE PRECONSOR ATP-DEPENDENT CLP PROTEASE PROTECLY TIC SUBUNIT (EC 3.4.21.92) ATP-DEPENDENT CLP PROTEASE PROTECLY TIC SUBUNIT (EC 3.4.21.92) CLPB PROTEIN A TP-DEPENDENT CLP PROTEASE ATP-BINDING SUBUNIT CLPX	A if Joseph Control of the Professes (FC 3 4.21.) Hypothetical Secretory Serine Protesse (FC 3 4.21.) ATP dependent Zn professes	XAA.PRO DIPEPTIDASE (EC 3.4.13.9) GAMMA.CI.UTAMYL.FRANSPEPTIDASE (EC 2.3.2.2) GAMMA.CI.UTAMYL.FRANSPEPTIDASE (EC 2.3.2.2) PENICILLIN.BINDING PROTEIN 5' PRECURSOR (O.ALANYL. D.ALANINE CARBOXYPEPTIDASE) (EC 3.4.11.9) XAA.PRO AMINOPEPTIDASE (EC 3.4.11.9)	PEPTION. DIPEPTIDASE DCP (EC. 34.15.5) AMINOPEPTIDASE N (EC. 3.4.11.2) AMINOPEPTIDASE N (EC. 3.4.11.2) VACUOLAR AMINOPEPTIDASE I PRECURSOR (EC. 3.4.11.1) XAA-PRO AMINOPEPTIDASE (EC. 3.4.11.9)	AMINOPEPTIDASE PROLYL ENDOPEPTIDASE (EC 3 4 21 26) AMINOPEPTIDASE GAMMA GLUTAMYLTRANSPEPTIDASE (EC 2 3 2 2) AMINOPEPTIDASE N (EC 3 4 11 2) PTRB periplasmic prolease	P FRB periplasmic protease (L42758) proteinase (Sueptomyces fividans) (L42758) proteinase (Streptomyces fividans) HEIC PROTEIN (EC 3 4 · · ·)	HFLC PROTEIN (EC 3 4 ··) O-SIALGGLYCOPROTEIN ENDOPEPTIDASE (EC 3 4 24 57) O-SIALGGLYCOPROTEIN ENDOPEPTIDASE (EC 3 4 24 57) O-SIALGGLYCOPROTEIN ENDOPEPTIDASE (EC 3 4 24 57)
Gene Name	EC-clpP, BS-clpP EC-clpB	EC-clpX, BS-clpX	BS.yvyE,EC.yıgZ BS.yqhT,EC.b2385	ECda	BS.yuiE,EC.pepA	EC-puB	EC-104-05 BS-ydie EC-1910
Stop	2497 137 798 3920	1072 9781 4453 5	507 117 507 121 1826	959 1067 3152 94	957 50 550 3933 1580	6193 2660 4949	5880 3965 1643 2149 3187
Start	3687 742 1388 2205	2349 10722 3620 862	5664 767 1 846 738	2289 1353	1738 1738 125 3430 207	4075 4778 1647 5194	7175 4919 969 1643 2156
Contig	GR00016 GR00152 GR00152 GR00464	GR00310 GR00202 GR00228 GR00324	GR00665 GR00751 GR00801 GR10005 GR00022	GR00125 GR00242 GR00289 GR00290 GR00323	GR00329 GR00337 GR00368 GR00548 GR00589 GR00589	GR00163 GR00275 GR00276	GR00023 GR00731 GR00125 GR00125
Identification Code	RXA00112 RXA00566 RXA00587	KXAU1999 RXA01120 RXA00744 RXA00844	RXA02317 RXA02644 RXA02820 RXA02859	EXA00499 EXA00877 EXA01014 EXA01018 EXA01147	RXA01161 RXA01181 RXA01277 RXA01914 RXA02000 RXA02000	RXA00621 RXA00622 RXA00977 RXA00982	RXA00152 RXA02558 RXA00500 RXA00501 RXA00501

### Enzymes in general

Function	ALPHA-RIBAZOLE-5'-PHOSPHATE PHOSPHATASE (EC 3 1 3 -) 3-OXOADIPATE ENOL-LACTONASE (EC 3 1 1 24) 4-AMINOBUTYRATE AMINOTRANSFERASE (EC 2 6 1 19) BETA C-5. LYASE (EC 3 · · · · ) PUTATIVE AMINOTRANSFERASE GLYCOSYL TRANSFERASE Acetylitansferases Acetylitansferases	Acetyliransferases Acetyliransferases Acetyliransferases (the isoleucine patch superfamily) Predicted methyliransferases Predicted S.adenosylmethionine-dependent methyliransferase SAM-dependent methyliransferases SAM-dependent methyliransferases SAM-dependent methyliransferases MODIFICATION METHYLASE (EC 2 1.173)	LACCASE I PRECURSOR (EC 1 10 3 2) LACCASE I PRECURSOR (EC 1 10 3 2) CARBONIC ANHYDRASE (EC 4 2.1 1) THIOL PEROXIDASE (EC 1 11 1 -) 2-NITROPROPANE DIOXYGENASE (EC 1 13 11 32)	Hypothetical Oxidoreduclase Hypothetical Oxidoreduclase Hypothetical Oxidoreduclase BETAINE ALDEHYDE DEHYDROGENASE PRECURSOR (EC 1 2 1 8) MORPHINE 8 DEHYDROGENASE (EC 1 1 1 218) NITRILOTRIACETATE MONOOXYGENASE COMPONENT A (EC 1 14 SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (MADP+) (EC 1 2 SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (MADP+) (EC 1 2 SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (MADP+) (EC 1 2	SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (NADPH) (EC 1 2 SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (NADPH) (EC 1 2 SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (NADPH) (EC 1 4 99 1) D-AMINO ACID DEHYDROGENASE LARGE SUBUNIT (EC 1 4 99 1) D-AMINO ACID DEHYDROGENASE LARGE SUBUNIT (EC 1 4 99 1) D-AMINO ACID DEHYDROGENASE SMALL SUBUNIT (EC 1 4 99 1) D-AMINO ACID DEHYDROGENASE SMALL SUBUNIT (EC 1 4 99 1) NADPH DEHYDROGENASE SMALL SUBUNIT (EC 1 4 99 1) NADPH DEHYDROGENASE SMALL SUBUNIT (EC 1 4 99 1) Oxoglutarate semialdehyde dehydrogenase (EC 1 2 1-)
Gene Name	BS-yonG,EC-gabf BS-palB,EC-malY	BS-dhbF,EC-eniF EC-yral,BS-yabC EC-yggH	EC.yacK EC.tpt,BS.ytgl BS.ytyB	BS-ydnJ	EC-gabD.BS-ycnH BS-ywmD
NT	648 1551 9812 1478 17703 449	1562 17387 3213 257 13040 26012 2189 17707	3130 5 11201 6 5208	971 1936 775 523 1070 1758	1228 8450 3160 5 4 4 4720 5053 602 1749
NT	1355 789 11155 2452 16561	16827 7034 1102 13804 26838 1589 18477	1640 592 10581 374 4186	1363 1226 1401 2 132 2544	608 9439 1538 598 831 7548 4821 5852 1573
Contig	GR00692 GR00308 GR00389 GR00489 GR00639	GR00245 GR00250 GR00758 GR00424 GR00741 GR0032 GR00519 GR00519	GR00351 GR00354 GR00715 GR00225	GR00337 GR00126 GR00180 GR00643 GR00679 GR00679	GR00170 GR00389 GR00209 GR00210 GR00296 GR00296 GR00296 GR00296
Identification	RXA02384 RXA01115 RXA01341 RXA01728 RXA02148 RXA02762	RXA00897 RXA02214 RXA02216 RXA01489 RXA01257 RXA02589 RXA00226 RXA01885 RXA02592	RXA01214 RXA01250 RXA02477 RXA02477		RXA0059 RXA01140 RXA011498 RXA00791 RXA01057 RXA01055 RXA01056 RXA01056

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				. ,		14), hippurate hydrolase												,				•	•		
		5	N-ACYL L-AMINO ACID AMIDOHYDROLASE (EC 35 1 14)	. 2 (	N-ACYL L. AMINO ACID AMIDOHYDROLASE (EC 3 5 1 14)	N-ACYL L-AMINO ACID AMIDOHYDROLASE (EC 3.3.1.14)		Hypothetical Melal Dependent Hydrolase	Predicted hydrolases (denalogenase related)	Predicted nyarotases (rons superioring)	Oradinad Znidenendent hydrolases	SALICYLATE HYDROXYLASE (EC 1 14 13 1)	SOLUBLE EPOXIDE HYDROLASE (SEH) (EC 3 3 2 3)	ACETYL HYDROLASE (EC 3.11)	PUTATIVE SECRETED HYDROGASE	SIALIDASE PRECURSOR (EC 3.2.1.18)	SIALIDASE PRECURSOR (EC 3 2 1 18)	SIALIDASE PRECURSOR (EC 3 2 1 18)	FN7VMF	METAL-ACTIVATED PTRIDOANL CITETING		2 NITROPROPANE DIOXYGENASE (EC 1 13.11.32)	ALCOHOL DEHYDROGENASE (EC 1.1.1.1)		the motabolism of inordanic compounds
	Gene Name		- <sup>-</sup>	-0			BS.hp0	EC-5130/ BS-yabD,EC-ycfH		EC-20844	BS-ykqC	BS-yhll	ac.whu		EC-51107.8S-ybbD							1	65-yrps EC-b0025		motaholism of it
z	Slop		694	1265	182	740	2877	5042 LTI	1673	4291	4489	127	35	1922	1840		4	1300	B24	1951	S	4346	5208		041.00
N	Start	ĺ	2 0ra	3 = 3	3104		1693	3657	2302	3461	3555	792	7	056 5	25.5	3	1200	1719	ድ	. 100	637	4035	4186	000	
	Config.		- GR00247	GR00247	GR00631	- GR00773	GR007 34 GR 10002	GR00003	GR00569	CROWING CRODS09	GR00236	GR00703	GR00287	GR00016	GR00555	ec /0MD	CR00278	GR00278	GR00722	10000	GR00167	GR00304	CR00354	GR00438	
Ldon'iffration	Code		RXA00905	PXA00906	RXA0030/ RXA02101	RXA02565	RXA02567	EXA00026	RXA01971	RXA00354	FXA01802	HXA00800	RXA00961	RXA00111	RXA01932	RXA02574		CXA00363	EXA02513		RXA00636	RXA00903	RXA01224	RXA01571	

# Genes encoding enzymes for the metabolism of inorganic compount

	Function	NITRATE REDUCTASE ALPHA CHAIN (EC 17994) NITRATE REDUCTASE ALPHA CHAIN (EC 17994) NITRATE REDUCTASE ALPHA CHAIN (EC 17994) NITRATE REDUCTASE ALPHA CHAIN (EC 17994)	
· .•		*	
	Gene Name	BS.narG,EC.narG	
	Stop		
	Start	370 686 1211	:
Ĕ	Config	GR00378 GR00377 GR00378	2000
N-metabolism	Idenlification	RXA01302 RXA01307 RXA01308	RXA01309

Function	NITRATE REDUCTASE ALPHA CHAIN (EC 1 7.994)	NITRATE REDUCTASE GAMMA CHAIN (EC 1.7.99.4)	NITRATE/NITRITE RESPONSE REGULATOR PROTEIN NARL	NITRATE/NITRITE RESPONSE REGULATOR PROTEIN NARP	NITRA TEANITRITE RESPONSE REGULATOR PROTEIN NARP	NITRATEINITRITE RESPONSE REGULATOR PROTEIN NARP	NITRATEINITRITE RESPONSE REGULATOR PROTEIN NARP	NITRALENITRIE SENSOR PHOLEIN NARK (EC. 2.7.3.)	Gene required for initiate assimination and anadropic grown	NUTIFICATION SUBSTANCE PROTEIN B	NH(3) DEPENDENT NAD(+) SYNTHETASE (EC 6 3 5 1)	NITRITE EXTRUSION PROTEIN	NITROGEN FIXATION PROTEIN FIXI (PROBABLE EI EZ TYPE CATION ATPASE) (EC 361.)	NITROGEN REGULATION PROTEIN NIFR3	NITROGEN REGULATORY PROTEIN P.II	NODULATION ATP-BINDING PROTEIN I	NODULATION PROTEIN N	OXYGEN-INSENSITIVE NAO(P)H NITROREOUCTASE (EC 1 · · · · )	PLASMID PTOM9 FROM ALCALIDENES XYLOSOXIDANS NREA AND NREB GENES, COMPLETE COS			Function	UREASE ALPHA SUBUNIT (EC 3.5.1.5) UREASE ALPHA SUBUNIT (EC 3.5.1.5) UREASE GAMMA SUBUNIT (EC 3.5.1.5) UREASE OPERON URED PROTEIN	UNEASE ACCESSORY PROTEIN UREF UREASE ACCESSORY PROTEIN UREG Uneaselhydrogenase-associated predicted GTPases	
Gene Name	i i	8S-narl,EC-narl				BS-yocG, EC-uhpA			BS ABIA, EC MORA	BS-nusA,EC-nusA	BS-nadE,EC-nadE	EC.nav		BS-yarF,EC-yhdG	EC glnX	BS-yviR		EC-51008	BS·yvgZ			Gene Name	BS-ureC BS-ureA	EC-y9(D	
Stop	1048	560 260	3686	1013	3382	1937	152	2951	/61	1937	2104	390	417	4350	797	14472	1369	9390	3741		K	Stop	4 1604 153 4268	2102 2782 3416 1868	
NT	1521	901	2997	102	4017	2545	123	1752	1033	2932	1274	1724	620	3208	-	15350	1001	8782	3442		Z	Slart	123 3 452 3420	1632 · 2105 2802 2734	
Contig	GR00810	GR00610 GR00610	GR00119	GR00021	GR00169	GR00339	GR00449	GR00119	, GR00444	GR00203	GROOM	GR00376	GR00412	GR00205	GR00764	GR00763	GR00221	GR00296	CR00385	-		Config	GR00655 GR00656 GR00655 GR00656	GR00656 GR00656 GR00656 GR00650	
Identification Code	RXA02017	RXA02018	RXA00471	RXA00133	RXA00650	RXA01189	RXA01607	RXA00470	RXA01589	RXA00756	BYA01073	COLLONA	RXA01412	RXA00773	RXA02748	RXA02745	RXA00820	RXA01059	RXA01324	Urease	Identification	Code	RXA02264 RXA02274 RXA02265 RXA02278	RXA02275 RXA02276 RXA02277 RXA02215	

## Phosphate and Phosphonate metabolism

	Gene Name	EC.phuA				BS-ytaK	BS.phoH, EC.bubbu	EC-pta, BS-pta	EC.pslS					1	BS-phoD	EC-ppa	•	8S-ykoX	•	EC-dedA	
¥	Slop	1783	5962	4	2044	. 177	15341	.2550	8246	1903	1083	₹	10985	1199	91001	14896	2292	101	2774	525	4260
Ä	Slart	2124	6375	294	1772	1222	14325	3932	6206	158	1467	384	10059	5193	8469	15169	1426	9512	3355	62	205
. •	Config	GR00638	GR00012	GR00632	GR00248	GR00173	GR00242	GR00418	CR00205	GR00491	GR 10016	CR00486	GR00720	GR00029	GR00422	GR00424	GR00447	CR00014	GR00162	_ CR00602	CR00636
Idenlification	Code	81100000	BXA00078	PXA02105	0×400912	EXA00663	PXADDBB	OXAD1437	DXA00778	PXA01732	BXA02877	BXA01716	DXA02497	DYA00191	BX401477	BXA01509	BXA01593	BXA00100	RXA00615	BXA02010	RXA02120

### Sulfate metabolism

Gene Name	BS-ygcA,EC-b0935
Stop	6 293 2644 733
Slart	448 3 1469 161
Config.	GR00012 GR00727 GR00211 GR00342
Identification	RXA00072 RXA02548 RXA00793 RXA001192

### Function

CARBOXYVINYL CARBOXYPHOSPHONATE PHOSPHORYLMUTASE (EC 2 7 8.23) PHOSPHONATES BINDING PERIPLASMIC PROTEIN PRECURSOR PHOSPHATE BINDING PERIPLASMIC PROTEIN PRECURSOR PHOSPHOPANTETHIENE PROTEIN TRANSFERASE, PPT IP ALKALINE PHOSPHATASE D PRECURSOR (EC 3 | 3.1) PHOSPHATE ACETYLTRANSFERASE (EC 2.3 1 8) INDRGANIC PYROPHOSPHAFASE (EC 3.6.1.1) 4-NITROPHENYLPHOSPHATASE (EC 3 1.341) DEDA PROTEIN, similar to alkaline phosphatase EXOPOLYPHOSPHATASE (EC 3 6 1 11) EXOPOLYPHOSPHATASE (EC 3 6.1.11) ALKALINE PHOSPHATASE (EC 3.1 3 1) PHOH PROTEIN HOMOLOG PHOH PROTEIN HOMOLOG PHAIA, B, C, D, E, F. OI GENES DEDA PROTEIN DEDA PROTEIN PHNB PROTEIN PHNB PROTEIN PHNA PROTEIN

### Function

PHOSPHOADENOSINE PHOSPHOSULFATE REDUCTASE (EC 1 8 99 4) SULFATE ADENYLATE TRANSFERASE SUBUNIT 2 (EC 2 7 7 4) SULFATE STARVATION INDUCED PROTEIN 8 SULFATE STARVATION INDUCED PROTEIN 6

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Function	SULFITE OXIDASE (EC 1.8.3.1) THIOSULFATE SULFURTRANSFERASE (EC 2.8.1.1) THIOSULFATE SULFURTRANSFERASE (EC 2.8.1.1) THIOSULFATE SULFURTRANSFERASE (EC 2.8.1.1)	ADENYLYLSULFATE KINASE (EC 2 7.1.23)		Function	FERRIC EN FEROCHELIN ES FERASE HOMOLOG FERRIC UPTAKE REGULATION PROTEIN FERRIPYOCHELIN BINDING PROTEIN FERRITIN	HEMIN BINDING PERIPLASMIC PROJEIN TRICE TO THE STATE OF T	IRON(III) DICITRATE BINDING PERIPLASMIC PROTEIN PRECURSOR IRON(III) DICITRATE BINDING PERIPLASMIC PROTEIN PRECURSOR IRON(III) DICITRATE BINDING PERIPLASMIC PROTEIN PRECURSOR	IRON(III) DICITICATE BINDING PERIPLASMIC PROTEIN PRECURSOR IRON(III) DICITICATE BINDING PERIPLASMIC PROTEIN PRECURSOR PERIPLASMIC IRON-BINDING PROTEIN SHIB	FERRIC ANGUIBACTIN-BINDING PROTEIN PRECUNSOR
Gene Name	EC-sseA	EC cysN		Gene Name	EC-fur,BS-yqN EC-b3279,BS-ytoA		EC.fec8 BS.yvrC	BS-yliY EC-bcp	,
NT	2497 2914 485	7884		Stop	706 3887 7749	258	827 2370 1241 1757	3532 3795 380	5402
Slart	1811 2120 1306	8837		Start	1848 3436 7192 548	980	1486 3287 2185 2892	2585 4586 3	4389
Config.	GR00356 GR00188 GR00463	GR00037	olism	Config.	GR00567 GR00011 GR00555	GR00586 GR00511 GR00118	GR00302 GR00013 GR00358	GR00451 GR00624 GR10016	GR00078 GR00013
Identification Code	. '	RXA02334 C	Fe-Metabolism	Identification	RXA01967 RXA00070 RXA01934	RXA01997 RXA01810 RXA00467	RXA01082 RXA00085 RXA01236	RXA01354 RXA01620 RXA02052 RXA02875	EXA00372 PXA00088
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Function	MAGNESIUM-CHELATASE SUBUNIT CHLI MAGNESIUM-CHELATASE SUBUNIT CHLI MAGNESIUM-CHELATASE SUBUNIT CHLI		
Gene Name			
Slop	. 789 1555		
Slad	570 1532 2004		
Config.	GR00474 GR00524 GR00524		
Identification	RXA01691 RXA01848 RXA01849		

# Modification and degradation of aromatic compounds

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	55.12)			CTASE COMPONENT (EC. 1.18.1.J)		14 (3 8)	14 13 8)	· r	ā	
Function	ARYLALCOHOL DEHYDROGENASE (NADP+) (EC 1 1.191) J.CARBOXY, CIS, CIS, MUCONATE CYCLOISOMERASE (EC 5.5.1.2) J.CARBOXY, CIS, CIS, INUCONATE CYCLOISOMERASE HOMOLOG (EC 5.5.1.2)	4.CARBOXYMUCONOLACTONE DECARBOXYLASE (EC 4 1 1 44) 4.CARBOXYMUCONOLACTONE DECARBOXYLASE (EC 4 1 1 44) MUCONATE CYCLOISOMERASE (EG 5 5 1 1) MUCONATE CYCLOISOMERASE (EG 5 5 1 1)	MUCONOLACTONE ISOMERASE (EC 5.3.3.4) 4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1.4) 4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1.4) 4-HYDROXYBENZOATE OCTAPRENYLTRANSFERASE (EC 2.5.1.4)	BENZALDEHTDE DEHTONO DEN STATEM FERREDOXIN – NAD(H) REDUCTASE COMPONENT (EC BENZENE 1,2-DIOXYGENASE SYSTEM FERREDOXIN – NAD(H) REDUCTASE COMPONENT (EC BENZENE 1,2-DIOX 1,2-DIOX 1,2-DIOXYGENASE III (EC 1,13-11,39)	CAFFEOTI CON CHIEFFILM SFERASE (EC 2 1 1 104) CAFFEOTI CO O WETHYLRANSFERASE (EC 2 1 1 104) CATECHOL 1.2 104 VGENASE (EC 1 13 11.1) CATECHOL 1.2 105 PHENE DESULFURIZATION ENZYME A	DIBENZOTHIOPHENE DESULFURIZATION ENZYME A DIBENZOTHIOPHENE DESULFURIZATION ENZYME C DIBENZOTHIOPHENE DESULFURIZATION ENZYME C DIBENZOTHIOPHENE DESULFURIZATION ENZYME C	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 2 (EC. 1.14.13.8) PARANITAOBENIZYL ESTERASE (EC. 3.1.1.3) PARANITAOBENIZYL ESTERASE (EC. 3.1.1.3)	PHENOL 2 MONOOXYGENASE (EC 1 14 13 7)	PROTOCATECHUATE 14-DIOXYGENASE BETA CHAIN (EC 1.13.11- PROTOCATECHUATE 13-DIOXYGENASE ALPHA SUBUNIT (EC 1.14.12-) FOLINATE 1, DIOXYGENASE ALPHA SUBUNIT (EC 1.14.12-)	TOLUATE 1,2 DIOXYGENASE BETA SUBUNIT (EC 1 14 12.)
Gene Name	EC 50419,BS yccK	ВЅ-ук <b>Ю</b>	F.C.ubiA	BS.//mT EC.b2542,BS.nasD BS.yfiE				BS-pribA		
NT Stop	1882	•	8025 2655 8737 ·					٦,	w, v,	2573 3107
Start	938	651 1098 1556 4121	9038 2945 7742	15614 8385 15	4644 5223 865	1909	1720	321	463 1167 1083	1533
Config	GR00003 GR00725	GR00794 GR00307 GR00637 GR00421								
Identification	EXA00024	RX402813 RX402113 RX402126	RXA02316 RXA01464 RXA02603	RXA02839 RXA02674 RXA01502	RXA02828 RXA01918 RXA02064	RXA00639 RXA00797 RXA01653	RXA00792 RXA02530 RXA02083	RXA00892 RXA02092 RXA00658	PXA01385 PXA01461 RXA01462	RXA00641 RXA00641 RXA00642

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Function	TOLUATE 1,2-DIOXYGENASE ELECTRON TRANSFER COMPONENT VANILLATE DEMETHYLASE (EC 1 14 ···) VANILLATE DEMETHYLASE OXIDOREDUCTASE (EC 1 ···) VANILLATE DEMETHYLASE (EC 1 1 1 ···)  XYLENE MONGOXYGENASE ELECTRON TRANSFER COMPONENT  1-hydroxy-2-naphihoale 1,2-dioxygenase (EC 1 ···) 1 ···)  CHLOROCATECHOL 1,2-DIOXYGENASE  CHLOROCATECHOL 1,2-DIOXYGENASE  QUINOLINATE SYNTHETASE A  GUINOLINATE SYNTHETASE A  GUINOLINATE SYNTHETASE (EC 1 ···) 1 ···)  MALEYLACETATE REDUCTASE (EC 1 ···) 1 ···)  A)-HYDROXYPHENYL, PROPIONATE HYDROXYLASE  POSSIBLE 2-HYDROXYHEPTA-2,4-DIENE-1,7-DIOATE ISOMAERASE  POSSIBLE 2-HYDROXYHEPTA-2,4-DIENE-1,7-DIOATE ISOMAERASE	SUCCINYL.COA 1-KETOACID-COENZYME A TRANSFERASE PRECURSOR (CC 2 8 3 3) SUCCINYL.COA COENZYME A TRANSFERASE (EC 2 8 3 3) SUCCINYL.COA COENZYME A TRANSFERASE (EC 2 8 3 3) 3.0XOADIPATE ENDL.LACTONASE (EC 3.1.1.24) CYTOCHROME P450 118 (EC 1.141
		F ( +
Gene Name	EC.h1803  BS.nadA  EC.fucO  EC.b0347,BSyhjG  BS.vieK,EC.b1180	85.yap.cc.stoD 85.yafD.EC.stoD EC.42920
NT Stop	4657 386 670 11347 7753 451 5 6 2360 1188 5593 5484 290 1428	336 973 1210 1644 1551 7269
Slart	3122 1 2 373 6589 6589 1575 826 671 1458 304 4310 4657 3 340 2185	949 1713 2715 2018 789 7751
Conlig	GR00168 GR00584 GR00584 GR00726 GR00726 GR00710 GR00710 GR00018 GR00018 GR000710	GR00309 GR00309 GR00305 GR00372 GR00308 GR00179
Identification Code	RXA00643 RXA01993 RXA01993 RXA01994 RXA02535 RXA01466 RXA021496 RXA02149 RXA00178 RXA00177 RXA00177 RXA00178	RXA01117 RXA0172 RXA01288 RXA01288 RXA01115

# Modification and degradation of aliphatic compounds

Function	ALKANAL MONOOXYGENASE ALPHA CHAIN (EC 1 14.14.3) ALKANAL MONOOXYGENASE ALPHA CHAIN (EC 1 14.14.3) ALKANAL MONOOXYGENASE ALPHA CHAIN (EC 1.14.14.3)	ALKANAL MONOOTOEMASE ACTION OF THE STATE OF
Gene Name	BS.yvbT,EC.yhbW	BS.ylnJ
Stop	6633 15385 820	560 7192 1070
Start	7376 16086 2	1603 6590 132
Contig	GR00048 GR00057	GR00750 GR00555 GR00555
Idenlification	FXA00299 FXA00332	RXA01838 RXA02643 RXA01933 RXA02351

# TABLE 2: GENES IDENTIFIED FROM GENBANK

Reference		of pyruvate carboxylass corboxylase, recombinant DNA carrying said fragment, strains carrying the	strains," Patent: EP 0358940-A 3 03/21/90		micro-organisms with delegulated infedime deligibles, a constant of the second of 19442. A \$ 07/20/93		182 on the many characterization of the fish	gene from coryneform bacteria," Biochem. Biophys. Res Commun.	236(2):383-388 (1997)	Biotechnol, 51(2):223-228 (1999)	Kimuia, E. el al. "Molecular cloning of a novel gene, disk, which restrict the defendant derived from Brevibacles ium	laciosermentum," Bioici Biotechnol Biochem, 60(10), 1565-1570 (1996)		masc		2 oxoglutarate aminotranslerase small subunits		Ut.	in; aminoglycoside	Ie-5.semialdehyde		chase		te synthetase	notyttansfeldse	c dehydialase
Gene Punction		Phosphoenol pyruvate c		Thiconine deliydiatase								·		D.glutamate racemase	transketolase	Glutamine 2-oxoglutarat	aconitase	Replication protein	Replication protein; animoglycoside	N-acelylglutamate-5-semialdehyde	dchydrogenasc	Glutamine synthetase	cyclast	Argininosuccinate synthetase	Ornithine carbanolydiansiciase	3.dehydroquinate dehydialase
Gene Name		phB						murC, fisQ; fisZ		murC; flsQ	disR		disR1: disR2	murl	3	gliB, gliD	Sch	CD	rcp; aad	amC		glnA	hısF	angG	argF	Core
<u> </u>	Accession No.	A09073		4.466.70	A45581,	A45583,	A45585	AB003132		AB015023	AB018530		4 0010471	AB070624	771170BA	AB024708	A DA 2 647 A	AB027714	AB027715	A F 00 \$ 2 4 2	AL 1000 LA	AF005635	AF030405	AF030520	AF031518	C10340.14

Reference		Michmeier 1 et al "The role of the Corynebacterium glutamicum tel gene in	(p)ppGpp metabolism." Microbiology, 144.1853-1862 (1998)															op o	Jish Phoning hinsynthetic	Park, S. et al. "Isolation and analysis of metry, a metricular and analysis of metry, a metry, a metry, and metry, and metry a	encoding homoseline artifyllansitiast in Co.) iccom.						Daniel, N. a. a. "Ryangerian of the Corynchacterium glutamicum panD gene		overproduction in Escherichia coli," Appl. Environ micronim, 55(4), 550
Gene Function		lythivate carboxylasc	Dipeptide binding piotein; addinice phosphoribosyllians ferase; GTP	pyrophosphokinase	Arginine repressor	Inositol monophosphate phosphatase	Argininosuccinale lyasc	N. acetylelufanylphosphate teductase,	omithine acetyltransferase; N.	acetylglutamate kinase, acetylomithine	nansminase; ornithine	carbamoyltransferase; arginine repressor;	l argininosuccinate synthase;	argininosuccinate lyase	Enoyl, acyl carrier protein reductase	ATP phosphoribosylhansferase	Phosphoribosylformimino-5-amino-1-	phosphoribosyl-d-imidazolecaiboxamide	isomerase	Homoserine O acetyltransferase		D. L. J. connects emphasize	Denygloquinale, sy innousay	Glittamine amidolialisterase	nospinoticosyty in a conception of the conceptio	5-enolpyruvylshikimate 3-phosphate	synthusc	Laspartaic-alpha-decaiboxylase piecuisor	`
Gene Name		pyc	dciAE; apt; rel		artiR	imnA	unpo.	algn	alge, alg., alge.	argo, argi, argin	12912				Λ 4α.	Parity	Delit	NSA.		melA			аюВ	hisH	hisE	aroA		panD	
GenBankin	Accession No.	AF038548	AF038651		AE041436	AL041430	AF045998	AF048764	AF049897							AF050109	A1-050100	AF051846		4 1705767	A1 02022		AF053071	AF060558	AF086704	AF114233		AF116184	

Accession No. AF124518 AF124600 AF124600 AF145897 AF145898 AJ001436 AJ004934 AJ007732 AJ010319 AJ132968 AJ132968	Gene Name aroD; aroE aroC; aroK; aroB; pepQ inhA inhA ectP dapD dapD fix, glnB, glnD, srp; amiP cal miqo	Gene Function  3-dehydroquinase; shikimate dehydrogenase Chorismate synthase; shikimate kinase; 3- dehydroquinate synthase; putative cytoplasmic peptidase  Transpun of ectoine, glycine betaine, proline Tetrahydrodipicolinate succinylase (incomplete') Phosphoenolpynvate-carboxylase; 9; high affinity ammonium uptake protein; putative omithine-cyclodecarboxylase; satcosine oxidase Involved in cell division; PH protein; uridylyfransferase (uridylyl-temoving enzmye); signal recognition particle; low affinity ammonium uptake protein Chloramphenicol aceteyl transferase Lenafate: quinone oxidoreductase	Peter, H. et al. "Corynebacterium glutamicum is equipped with four secondary carriers for compatible solutes. Identification, sequencing, and characterization of the proline/ectione uplake system, ProP, and the ectione/proline/glycine betaine carrier, EctP," J Bacteriol, 180(22):6003-6012 (1988) Wehrmann, A. et al. "Different modes of diaminopimelate syndhesis and their role in cell wall integrity. A study with Corynebacterium glutamicun," J Bacteriol, 180(12):3159-3165 (1998)  Jakoby, M. et al. "Nitrogen regulation in Corynebacterium glutamicum," Isolation of genes involved in biochemical characterization of corresponding proteins," FEMS Microbtol, 173(2):303-310 (1999)  Molenaar, D. et al. "Biochemical and genetic characterization of the membrane-associated malate dehydrogenase (acceptor) from Corynebacterium glutamicum," Eur. J Biochem, 254(2):395-403 (1998)
A1238250 A1238703	hdn Piod	NADII dehydrogenusc Porin	Lichingei, T. et al "Biochemical and biophysical characterization of the cell wall porin of Corynchacterium glutamicum. The channel is formed by a low to a part of the cell o
D17429		Transposable element 1831831	molecular mass polypeptide, Biocitemistry, 3 (43), 13024-13034 (1770). Vertes, A.A. et al. "Isolation and characterization of 1831831, a transposable element from Corynebacterium glutamicum," Mol Microbiol, 11(4), 739-746 (1994).

		F	Reference
GenBank"	Gene Name	Cent Function	
Accession No. D84102	Vypo	2-oxoglutarate deliydiogenasc	Usuda, V. et al. "Molecular cloning of the Corynebacterium gintamicum (Brevibacterium lactofermentum AJ12036) odhA gene-encoding a novel type (Brevibacterium lactofermentum AJ12036) odhA gene-encoding a novel type
		Hangering delivideoenase: homosciine	of 2-oxogludarate denyoungulaxe, first consideration of 1-thereonine and 1-tsoleucine, Patent: JP Katsumata, R. et al. "Production of 1-thereonine and 1-tsoleucine," Patent: JP
E01358	hdh, hk	kinase	1987232392-A 1 10/12/87
E01359		Upstream of the start codon of homoserine kinase gene	1987232392-A 2 10/12/87
E01375		Тгургорны орстоп	Marini K et al "Tryntonhan oncton, peptide and protein coded thereby,
E01376	ருப்; ரிந்	Leader peptide; anthranilate synthase	utilization of tryptophan operon gene expression and production of
٠			Majeri K et al. "Tryptophan operon, peptide and protein coded thereby,
E01377		Promoter and operator regions of	utilization of tryptophan operon gene expression and production of tryptophan approach 1 10/24/87
		Biolin-synthase	Hatakeyama, K. et al "DNA fragment containing gene capabic of course
E03937	-		biolin synthelists and its contraction, when we have an inorparsfer and whoman ket at "Gene coding disminopelargonic acid an inorpansfer as and
E04040		Diamino pefaigonic acid aminoffailstefase	desthiobiotin synthetase and its utilization," Patent: Jp 1992330284-A 1
			11/18/92
E04041		Desthiobiotinsynthetasc	desthiobioin synthetase and its unlization," Patent 1P 1992330284-A 1
			11/18/92 "C - DNA coding ashardase and utilization thereof," Patent.
E04307		Flavum asparlase	Kurusu, Y. et al. Octic Dixix course aspect.  Jp 1993030977-A 1 02/09/93  Leave the controlling DNA." Patent JP
E04376		Isocitric acid lyasc	Katsumala, R. et al. Oche mainteatation controlline DNA " Patent. JP
E04377		Isocihic acid Iyase N-terninal fragment	Katsumata, R et al. "Gene maintestanon como como proposos 73 03/09/93
E04484		Prephenate dehydratase	Solouchi, N. et al. "Production of L-phenylauning by ichingment, N. et al. "Patent: JP
E05108		Aspartokinusc	Fugono, N. et al. "Gene DNA coung Aspartoknings and mg 1993184366-A 1 07/27/93
E05112		Diliydro-dipichorinate synthetase	Hatakeyama, K. ci al "Gene LINA Coung uniyu Gerrangana," Patent IP 1993184371-A 1 07/27/93

Accession No. E05776 E05779 E06110 E06111 E06825 E06825	Gene Name	Usaminopime lie acid deliydiogenase  Threonine synthase  Prephenate dehydratase  Mutated Prephenate deliydiatase  Acetohydroxy acid synthetase  Aspartokinase  Mutated aspartokinase alpha subunit  Mutated aspartokinase alpha subunit	Kubayashi, M. et al. "Gene DNA coding Diaminopimelic acid dehydrogenase and its use," Patent. JP 1993284970. A 1 11/02/93  Kohama, R. et al. "Gene DNA coding threonine synthase and its use;" Patent. JP 1993284972. A 1 11/02/93  Kikuchi, T. et al. "Production of L-phenylalanine by fermentation method," Patent. JP 1993344881. A 1 12/27/93  Kikuchi, T. et al. "Production of L-phenylalanine by fermentation method," Patent. JP 1993344881. A 1 12/27/93  Imui, M. et al. "Gene tapable of coding Acetohydroxy acid synthetase and its lmi, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 03/08/94  Sugimoto, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 03/08/94  Sugimoto, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 Sugimoto, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 03/08/94  Sugimoto, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 03/08/94  Sugimoto, M. et al. "Mutant aspartokinase gene," patent. JP 1994062866. A 1 03/08/94
E08177 E08177 E08178. E08179, E08180, E08181, E08182 E08233 E08234 E08234	secv	Aspartokinase  Feedback inhibition-released Aspartokinase  Acetohydroxy-acid isometoreductase  FT aminotransferase and desthiobiotin synthetase promoter region Biotin synthetase	protein to membrane," Patent: JP 1994169780. A 1 06/21/94  Salo, Y et al "Genetic DNA capable of coding Aspartokinase released from Salo, Y et al "Genetic DNA capable of coding Aspartokinase released from Salo, Y. et al. "Genetic DNA capable of coding Aspartokinase released from feedback inhibition and its utilization," Patent: JP 1994261766. A 1 09/20/94 feedback inhibition and its utilization," Patent: JP 1994261766. A 1 10/04/94  Asai, Y. et al. "Gene DNA coding acelohydroxy acid isomeroreductuse," Patent: JP 1994277073. A 1 10/04/94  Hatakeyama, K. et al. "DNA fragment having promoter function in Hatakeyama, K. et al. "DNA fragment having promoter function in Itatakeyama, K. et al. "DNA fragment having promoter function in sectorium," Patent: JP 1995031476. A 1 02/03/95

Con Rank 111	Gene Name	Gene Function	Reference
			K. cl. al. "DNA fraement having promoter function in coryneform
E08649		Aspartase	bacterium," Patent. JP 1995031478. A 1 02/03/95
E08900		Dihydrodipicolinate reductase	Madori, M. et al. "DNA Tragment Containing Bene County 2017 1 03/20/95 acid reductase and utilization thereof," Patent: JP 1995075578-A 1 03/20/95
E08901-		Diaminopimelic acid decarboxylase	Madoti, M. et al. "DNA fragment containing gene count transmining." Co. decarboxylase and utilization thereof," Patent. Jp 1995075579-A 1 03/20/95
E12594		Seine hydioxymethylliansferase	Hatakeyania, K. et al. "Production of L-trypophan, Falein of 1977-222371.
E12760,		transposase	Motiva, M. et al. "Amplification of gene using affiltent itemsposes, committee of 1997070291-A 03/18/97
E12758 E12764		Arginyl-IRNA synthelase; diaminopimelie	Moriya, M. et al. "Amplification of gene using artificial transposon," Patent.
E12767		acid decarboxylase Dihydrodipicolmic acid synthetase	Moniya, M. et al. "Amplification of gene using artificial transposon," Patent: Jp 1997070291-A 03/18/97
E12770		aspariokinasc	Moriya, M. et al. "Amplification of gene using artificial fransposon, Fatelli.  Jp 1997070291-A 03/18/97
E12773		Dihydrodipicolinic acid reductase	Moriya, M. et al. "Amplification of gene using artificial flansposon, Fateril 19 1997070291-A 03/18/97
E13655		Glucose 6-phosphate dehydrogenase	Halakeyama, K. et al. "Glucose-6-phosphate denydfolgenase and Divorgenase of coding the same," Patent: JP 1997224661-A 1 09/02/97
1.01508	IIvA	Threonine dehydralase	Mocckel, B. et al. "Functional and structural analysis of the tincommercation of Corynebacterium glutamicum," J. Bacteriol, 174,8065-8072
<u>L07603</u>	EC 4.2 1.15	3. deoxy. D. arabinoheptulosonate. 7. phosphate synthase	Chen, C et al. "The cloning and nucleotide sequence of Corymebacterium glutamicum 3-deoxy. D. arabinoheptulosonate. 7-phosphate synthase Bene", Fease Microbiol 120, 107,223,230 (1993)
L09232	IIvB; ilvN; ilvC	Acetohydroxy acid synthase large subunit; Acetohydroxy acid synthase small subunit;	Keilhauer, C. et al. "Isoleucine synthesis in Corynebacterium glutamicum. notecular analysis of the IIvB-iIvN-iIvC operon," J Bacteriol, 175(17).5595-5603 (1993)
		Attitution and assessment	

		Cone Renefion	Reference
GenBank <sup>12</sup>	Cene iname		all of the
Accession No.	PrsM	Phosphoenolpyruvate sugar	Fouel, A et al. "Bacillus subtilis sucrose-specific encyment of and homology to
		phosphotransicrase	enzymes 11 from enteric bacteria," PNAS USA, 84(24) 8773-8777 (1987); Lee,
		*	J.K. et al. "Nucleotide sequence of the gene encoding the Corynebacterium
			glutamicum mannose enzyme 11 and analyses of the deduced protein
			sequence," FEMS Microbiol Lett. 119(1-2).131-143 (1994)
L27123	Вхв	Malaic synthase	Lee, H. S. et al. "Molecular characterization of accu, a gene circums."  compasse in Cornebacterium glutomicum," J. Microbiol. Biotechnol.
			4(4) 256-263 (1994)
127126		Pymvalc kinase	Jetten, M. S. et al. "Structural and functional analysis of pytovals and Consumple of the Consumply of the C
			(1994)
1,202.60	Ace	Isocitrate tyasc	TAIL Sequence analysis and
1.35906	dixi	Diphtheria loxin repressor	Oguiza, J.A. et al. Moleculai clonnig, Divis superior and J.A. et al. Moleculai clonning of the Correctation of the Correctati
			[actofermentum," J. Bacteriol, 177(2):465-467 (1995)
M13774		Prephenale dehydratase	Follettie, M.T. et al. "Molecular cloning and nucleotide sequence of the Corynebacterium glutamicum phcA gene," J Bacteriol, 167:695-702 (1986)
M16175	SSIRNA		Park, Y-H et al. "Phylogenetic analysis of the corynetorm bacteria by 30 (RNA sequences," J. Bacteriol, 169:1801-1806 (1987)
	Jun	Anthranilate synthase, 5' end	Sano, K. et al "Structure and function of the trp operon control regions of
M10005	1	1	Brevibacterium factolermentum, a grutamit actus processes 52, 191-200 (1987)
M16664	IrpA	Tryptophan synthase, 3'end	Sano, K. et al. "Structure and function of the try operan control regions of Brevibacterium lactofermentum, a glutantic acid-producing bacterium," Gene,
			52.191.200 (1987)
M25819		Phosphoenolpyruvate catboxylase	O'Regan, M. et al. "Cloning and nucleoline sequence of ""  Phosphoenolpyravate carboxylase coding gene of Corynchacterium  Archinga "Come 17/2/231 (1989)
		Mild and Mild and Miller See	Roller, C. et al. "Gram-positive bacteria with a high DNA G+C content are
N485106		255 TIONA Bene machinin address	characterized by a common insertion within their 23S 1RNA genes, J. Gen. Adversabled, 138, 1167-1175 (1992)

ConRanbia	Gene Name	Gene Punction	Reference
Accession No.		1	are leading the DNA G+C content are
M85107,		23S IRNA gene insertion sequence	characterized by a common inscriton within their 23S 1RNA genes," J. Gen
-			Microbiol, 136, 1107-1113 (1372)
M89931	aecD: bmQ, yhbw	Beta C.S lyase, branched chain annino acid uptake carrier, hypothetical protein yhbw	lyase with alpha, beta-climunation activity that degrades aninoctly lysteine."
			Coryncbacterium glutanicum ATCC 13032 is directed by the bmQ gene
			product, Area includion, 193(2) 252 (1932)
829299	tтр	Leader gene (promoter)	hyperproducing shain of Corynebacterium glutamicum: identification of a
	,		mutation in the trp leader sequence," Appl. Environ Microbiol, 39(3), 191-199 (1993)
U11545	ιφD	Anthranilate phosphorosylnansferase	O'Gara, J.P. and Dunican, L.K. (1994) Complete nucleotide sequence of the Corynebacterium glutumicum ATCC 21850 tpD gene." Thesis, Microbiology
			Department, University College Galvay, Heland
U13922	cgilM; cgilR, cigilR	Putative type 11.5 cytosoine methyltransferase; putative type 11	Schaler, A. et al. "Cloning and characterization of a 2017 of 500
*		restriction endonuclease; putative type I of	13032 and analysis of its role in intergenent conjugation (vin Estimation)
	· .	type III restriction endonucleuse	Corynebacierium glutamicum cgllM gene encoding a 5-cytosine in an MerBC-deficient Escherichia coli strain," Gene, 203(2):95-101 (1997)
7707113	Vie		dri caraci
U31224	ppx		Ankii, S. et al. "Mutations in the Cotynebacterium giudamicumpromic
			178(15) 4412-4419 (1996)
U31225	proC	L proline: NADP+ 5.0x idoreductase	Ankri, S. et al "Mutations in the Corynebacterium glutamicumprofiles biosymthetic pathway. A natural bypass of the prod step," J Bacteriol.
			178(15):4412-4419 (1996)
U31230	obg; proB, unkdh	9;gamma glutanyl kinase;similar to D. isonier specific 2-hydroxyacid	Ankri, S. et al. "Mutations in the Corynebacterium grudamicumpromise bijosynthetic pathivay. A natural bypass of the proA step," J. Bacteriol.
		dehydrogenascs	(227) (11-7) (110/1

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Gen Bank <sup>n,</sup>	Gene Name	Gene Function	Reference
Accession No.	,		
U31281	bioB	Biolin synthasc	Screbriiskii, 1.G., "Two new members of the bio B superfamily: Cloning, sequencing and expression of bio B genes of Methylobacullus flagellatum and Corvebacterium plutamicum," Gene. 175-15-22 (1996)
U35023	thiR, accBC	Thiosulfate sulfurtransferase; acyl CoA carboxylase	Jager, W. et al. "A Corynebacterium glutamicum gene encoding a two-domain protein similar to biolin carboxylases and biolin-carboxyl-carrier proteins," Arch Microhiol, 166(2);76-82 (1996)
U4353S	СШІ	Multidrug resistance protein	Jager, W. et al. "A Corynchacterium glutamicum gene conferring multidrug resistance in the heterologous host Escherichia coli," J Bacteriol, 179(7):2449-2451 (1997)
U43536	clpB	Heat shock ATP-binding protein	
U53587	aphA-3	3'5". aminogly coside phosphotiansferase	
U89648		Corynebacterium glutanicum unidentified sequence involved in histidine biosynthesis, narial sequence	
X04960	ւթ∧; ւթΒ; ւթС; ւրD; ւթΕ; ւրG; ւր!	Tryptophan operon	Matsui, K et al. "Complete nucleotide and deduced amino acid sequences of the Brevibacterium factofermentum tryptophan operon," Nucleic Acids Res. 14(24):10113-10114 (1986)
X07563	lys A	DAP decarboxylase (meso diaminopiniclale decarboxylase, EC 4.1.1.20)	Yeh, P. et al. "Nucleic sequence of the lysA gene of Lotynebacterium glutanucum and possible mechanisms for modulation of its expression," Mal Gen Genea, 212(1):112-119 (1988)
X14234	EC 4 1.1.31	Phosphoenolpyruvate carboxylase	Eikmanns, B. J. et al. "The Phosphoenolpyrivate carboxylase gene of Corynebacterium glutamicum: Molecular cloning, nucleolide sequence, and expression," Mol Gen. Genet, 218(2):330-339 (1989); Lepiniec, I. et al. "Sorghum Phosphoenolpyrivate carboxylase gene family: structure, function and molecular evolution," Plant Mol Biol, 21 (3):487-502 (1993)
X17313	Ída	Fructose-bisphosphate aldolase	Von der Osten, C.H. et al. Motecular cioning, nucleorine sequence and structural snatysis of the Corynebacterium glutamicum fda gene: structural comparison of C. glutamicum fructose 1, 6 biphosphate aldolase to class 1 and class 11 aldolases," Mol Microbiol.
X53993	dapA	L.2, 3-dihydrodipicolinate synthetase (EC 4 2.1.52)	Bonnassic, S. et al "Nucleic sequence of the dapA gene 110m Corynebacterium glutamicum," Nucleic Acids Res., 18(21):6421 (1990)



		Cone Runction	Reference
GenBank <sup>1M</sup>	Cene Name		in the can bene encoding
X66078	copi	Ps1 protein	Jolist, G. et al. "Cloning and nucleolide sequence of the Co. et al. "Cloning and nucleolide sequence of the two major secreted proteins of Corynchacterium glutamicum:  PS1, one of the two major secreted proteins of Corynchaeterium antigen
			85 complex," Mol. Microbiol, 6(16),2349-2362 (1992)
X66112	118	Citrate synthase	Eikmanus, B.J. et al. "Cloning sequence, expression and analysis of the Corynebacterium glutanicum gltA gene encoding cittate synthase," Microbiol, 140,1817-1828 (1994)
X67737	dapB	Dihydrodipicolinate reductase	Borney 11 of all "Characterization of the LSDB gene encoding PS2, an ordered
X69103	csp2	Surface layer protein PS2	surface-layer protein in Corynebacterium glutamicum," Mol Microbiol.
			9(1)-97-109 (1993) Ronanty, C. et al. "Identification of 181206, a Corynebacterium glutamicum
X69104	-	ISJ related insertion cientein	183-related insertion sequence and phytogenetic analysis," Mol. Mictobiol.
	· · · · · · · · · · · · · · · · · · ·		14(2) 271:301 (1277)
X70959	lcuA	Isopropylnialate synthase	scrivities, structure of leud, and effect of leud inactivation on lysine scrivities, "Appl Environ Microbiol, 60(1), 133-140 (1994)
		12. Steele debydevocnace (NADP4)	Eikmanns, B.J. et al "Cloning sequence analysis, expression, and mactivalion
X71489	93	ISOCIII die deil deile d	of the Corynebacterium glutamicum ted gene encoding isocitrate dehydrogenase and biochemical characterization of the enzyme," J. Bacteriol,
			177(3).774.782 (1995)
770000	CDIIA	Glutamate dehydrogenase (NADI)+)	or a try from a try of only strain of
X75083,	mitiA	5-methyltryptophun resistance	Heery, D.M. et al. A sequence none in professional formation of the Concupied of the Concup
X70584			Biochem Biophys Res. Commun, 201(3)-1255-1262 (1994)
X75085	ICCA		Fitzpaurick, R. et al. "Construction and Linaracterium factofermentum," Applied Corynebacterium ghutamicum and Brewibacterium factofermentum, Applied Corynebacterium factofermentum, Applied Corynebacterium (Applied Applied
		11 - in the trade trace. 2	Reinscheid, D.J. et al. "Characterization of the 1socinate lyage gene from
X75504	aceA; thiX	l'aniai isocinaie iyax,	Corynchacterium glulamicum and biochemical analysis of the enzyme, J Bucleriol, 176(12):3474-3483 (1994)
X7687S		A TPase beta subunil	Ludwig, W et al. "Phylogenetic relationships of bacteria bases of a subunit sequence analysis of clongation factor. Tu and ATP-synthase beta subunit
			genes," Antonie Van Leeinvenhock, 64:283-305 (1993)



Conflantin	Gene Name	Gene Function	Kelerence
		יו	1 indivin W. cf al. "Phylogenetic relationships of bacteria based on comparanve
X77034	ja,	Elongation factor I u	sequence analysis of clongation factor Tu and ATP-synthase beta-subunit genes," Antonie Van Leeuwenhoek, 64.285-305 (1993)
	IECA		Billman Jacobe, H. "Nucleolide sequence of a teen general Corynebacterium glutamicum," DNA Seq. 4(6):403-404 (1994)
	яжВ	Malaic synthase	Reinscheid, D.J. et al. Walan Symmus. Programmer sequence analysis," pta-ack operon encoding phosphotransacetylase: sequence analysis,"  Microbiology, 140.3099-3108 (1994)
	16S rDNA	16S ribosomal RNA	Rainey, F.A. et al. "Phylogenetic analysis of the genus Norcardia Norcardia and evidence for the evolutionary origin of the genus Norcardia from within the radiation of Rhodococcus species," Microbiol, 141.523-528
	gluA; gluB; gluC, gluD	Glutamate uplake system	Kronemcyer, W. et al "Structure of the gluABCD cluster encoding the glutamate uptake system of Corynebacterium glutamicum," J Bucterial, 177(5):1152-1158 (1995)
	dapE	Succinyldiaminopimelate desuccinylase	Wehmann, A et al "Analysis of different DNA tragiliens of Corynebacterium glutanticum complementing dapE of Escherichia coli,"  Microbiology, 40:3349-56 (1994)
	16S IDNA	16S ribosomal RNA	Ruimy, R et al. "Phylogeny of me genus Chrystoles," Int. J. Syst. Bacteriol, analyses of small subinnit ribosomal DNA sequences," Int. J. Syst. Bacteriol, 45(4):740-746 (1995)
	asd; lysC	Asparlate semialdehyde dehydi ogenase: 9	Seichijski, J. et al. "Mullicopy suppression by assignments," J. dependent complementation by heterologous prod in prod mutants," J. Bucieriol, 177(24) 7255-7260 (1995)
	PioA	Gamma-glutamyl phosphate reductase	dependent complementation by heterologous prod in prod mutants," J  Bacteriol, 177(24) 7255-7260 (1995)
	16S IDNA	16S ribosomal RNA	Pascual, C. et al. Thylogenetic analysis of the Econor, 45(4):724.728 (1995) on 16S 1RNA gene sequences, 111. J. Syst Bacteriol, 45(4):724.728 (1995)
\	arop; dapE	Atomatic amino acid permease; ?	Wentmann, A. et al. Turkmonth and Corymebacterium glaamicumproline reveals the presence of anop, which cocodes the atomatic amino acid (tansporter," J. Bacteriol, 177(20), 5991. 5993 (1995)



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GenBank <sup>14</sup> G	Gene Name	(Sene Function	July work of the state of the s
Accession No.		Comment E45	Patek, M et al "Promoters from Corynchacterium glutamicum. crowns.
X90363		-	molecular analysis and search for a consensus month, which is a search for a consensus month, and the search for a consensus month, which is a search for a consensus month, and the search for a
		Elita Elita	Patek, M. et al. "Promuters from Corynebacterium glutamicum. cloning,
X90364		Promoter tragment or	molecular analysis and search for a consensus moun,
			Patek, M. et al. "Promoters from Corynebacterium glutamicum: cloning,
X90365		Promoter tragment 173	molecular analysis and search for a consensus moull, Millionings,
			142:1297-1309 (1996)
700366		Promotes fragment PF 101	Patek, Ni, et al. 1 millions of the consensus motif," Microbiology, molecular analysis and search for a consensus motif," Microbiology.
20004			142:1297-1309 (1996)
2,003,5		Promoter fragment PF 104	Paick, N. et al. Fromotics from Co. a consensus molif," Afici obtology, molecular analysis and search for a consensus molif," Afici obtology,
7905064			142.1297.1309 (1996)
0,000		Promoter fragment PF 109	Palek, M. et al. "Promoters from Coryncoartering" by moternia analysis and search for a consensus motif," After obiology.
80806X			142:1297-1309 (1996)
X93513	amt.	Animonium transport system	Siewe, R.M. et al. "I unctional and general general annuonium," J Biol Chem, uninonium uplake carrier of Corymebacterium glutamicum," J Biol Chem,
			271(10):3538-3403 (1970). Peter, 11 et al.: "Isolation, characterization, and expression of the
X93514	belP	Glycine betaine fransport system	Corynebacterium glutanticum belp gene, encoding the flansport system compatible solute glycine betaine," J Bacteriol, 178(17):5229-5234 (1996)
X95649	0rf4		dapA. ORF4 oper on of Corynchacterium glutamicum, encoding two enzynics dapA. ORF4 oper on of Corynchacterium glutamicum, encoding two enzynics
-		1 synorter molein 1 vsine export	involved in Llysing syndesis, Diorection, with a new type of cellular Viljic, M. et al. "A new type of transporter with a new type of cellular
X96471	lysE; lysG	regulator protein	function: L. lysine export from Colynepacterium Branch (22(5):813-826 (1996)

1		Gene Punction	Reference
GenBank	Cene Manne	6	in Coramphacterium plulamicum and
Accession No. X96580	panB, panC; xylB	3-niethyl-2-oxobulanoalc hydroxymcihyltransfcrase, pantoafe-bela- alanine ligase; xylulokinase	Sahm, H et al. "D pantolhenate synthesis in Colymbras for D pantothenate use of panBC and genes encoding L valine synthesis for D pantothenate overproduction," Appl Environ Microbiol, 65(5), 1973-1979 (1999)
		Insertion sequence 151207 and transposase	and acception of the pene encoding
X96962 X99289		Elongalion factor P	Ramos, A. et al. "Cloning, sequencing and expression clongation factor P in the amino-acid producer Brevibacterium lactofermentum clongation factor P in the amino-acid producer Brevibacterium glutamicum ATCC 13869)," Gene, 198:217-222 (1997) (Corynebacterium glutamicum ATCC 13869)," Gene, 198:217-222 (1997)
Y00140	thiB	Homoserine kinasc	Mateos, L.M. et al. "Nucleotide sequence of the homoserm." (1987) of the Brevibacterium lactofermentum," Nucleic Acids Res., 15(9):3922 (1987)
Y00151	qqy	Meso-diaminopimelate D.dehydiogenase (EC 14.1.16)	dehydrogenase gene from Corynebacterium glutamicum," Nucleuc Acids Res. 15(9):3917 (1987)
Y00476	thrA	Homoserine deliydrogenase	Mateos, L. M. et al. Nucleoting September 1. Nucleic Acids Res., (thi A) gene of the Brevibacterium factofermentum," Nucleic Acids Res., 15(24):10598 (1987)
Y00546	hom; thrf3	Homoserine dehydrogenase; homoserine kinase	Peoples, O.P. et al. Nucleonide sequence of Corymebacterium glutamicum hom-thr B operon," Mol Microbiol., 2(1):63-72 Corymebacterium glutamicum hom-thr acterization, and chromosomal
Y08964	murC, fisQ/divD; fisZ	UPD N. acetylmuraniale-alanine ligase, division initiation protein or cell division protein cell division protein	Honsubia, M. P. et al. Identification, Constraint factofermentum," Mol Gen organization of the fisz gene from Brevibacterium factofermentum, "Mol Genet, 259(1):97-104 (1998)
709163	puiP	High affinity proline transport system	glutanicumproline and characterization of a low-affinity uptake system for compatible solutes," Arch Microbiol., 168(2): 143-151 (1997)
γ09548	рус	Pyiuvate caiboxylase	glutamicum: characterization, expression and inactivation of the pyc gene,"  Microbiology, 144.915-927 (1998)
Y09578	leuB	3-isopiopylmalate dehydiogenase	Patek, M. et al. Analysis of the real Eur. (1998) glutamicum, Appl Microbiol, Biotechnol., 50(1):42-47 (1998)
Y12472		Attachment site bacteriophage Pht-10	construction of an integration vector," Adictobiol., 145:539-548 (1999)

		Paradion.	Reference
GenBanktu	Gene Name	Creme rubellon	
Accession No.	;		Deter 11 ct al "Corynchacterium glutamicum is equipped with four secondary
Y12537	prop	Proline/ectoine uptake system protein	carriers for compatible solutes. Identification, sequencing, and characterization
			of the prolinctectoine uptake system, Prop. and the ectoricyproline/Biyenic
,	. =		Ociame Callier, Lett., S. Barreller, S. Barreller, M. State Bloom of Corvnebacterium glutamicum gln & Bene
Y13221	glnA	Glutamine synthelase I	encoding glutamine synthetase 1," FEMS Microbiol Lett., 154(1):81-88 (1997)
V14647	lnd	Dihydrolipoamide dehydrogenase	. 1 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Y 18059		Attachment site Corynephage 304L	Moleau, S. et al. Analysis of the mic formages," Virology, 255(1) 150-159 (1999) integrase module among coryncphages," Virology, 255(1) 150-159 (1999)
721501	argS; lysA	Aiginyl-(RNA synthetase; diaminopimelate	Oguiza, J. A. et al. "A gene encoding arginyl-tikny symmetast is not accommendation of the loss gene in Brevibacterium lactofermentum.
		decarboxylase (partial)	Regulation of args-1/34 cluster expression by arginine," J
			Hacieriot, 113(22) 1330-1332 (1337)
221502	dapA; dapB	Dihydrodipicolinate synthase; dihydrodipicolinate reductase	Pisabano, A et al. A Chaice of thirs bereather the process, and a Bicvibacterium lactofermentum encodes dihydrodipicolinate reductase, and a hind polymentide of unknown function," J. Bacteriof, 175(9):2743-2749
			(1993)
229563	IlirC	Threonine synthase	Malumbres, M et al. "Analysis and expression of the fine feat of the three three synthase," Apul Environ Microbiol, 60(7)2209-2219 (1994)
	14C - DNA	Gene for 16S ribosomal RNA	Arreibacteriim
249822	sigA	SigA sigma factor	Oguiza, J A. et al "Mullipic signa factor genes in Dictinical (178(2):550-
			553 (1996) .: At 11DB or Jectors 4. Philpiciase 0.
249823	galE; dtxR	Catalytic activity UDP-galactose 4.	Oguiza, J A et al "The gall; gene encouing ine Our genevious of princing and Brevibacterium lactofennement is coupled transcriptionally to the dind?
		epinicrase; diplinicità tovili refinito)	gene," Gene, 177.103-107 (1996)
749824	orf!; sigB	?; SigB sigma factor	Oguiza, J A. et al "Multiple sigma factor genes in Dicyloacterior". Inacterior, 178(2):550-
			553 (1996)
7,665.34		Типьрозвяс	Concia, A. et al. "Clouing and chattecterization by all 13-1118, conserved the periodic of Breybaclerium factofermentum ATCC 13869," Gente,
			170(1) 91-94 (1996)
	,		the absolute the investigity of the firebour approximation of the same of the

TA sequence for this gene was published in the indicated reference. However, the sequence obtained by the inventors of the present application is significantly published wersion. It is believed that the published version relied on an incornect start codon, and thus represents only a fragment of the actual coding region.

TABLE 3: Corynebacterium and Brevibacterium Strains Which May be Used in the Practice of the Invention

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	Navum	21127						
Brevibacterium	Navum		B11474	14				
Brevibacterium	hcalii	15527					1	
Brevibacterium	keloglutamicum	21004						
Brevibacterium	ketoglutamicum	21089					-	
Bievibacterium	ketosoreductum	21914					1	
Brevibacterium	lactofermentum			70			1	
Brevibacierium	lactofemicntum	0		74				
Brevibacterium	lactofermentum			77				
Bievibacterium	lactofermentum	21798		•			1	
Brevibacterium	lactofernientum	21799						
Brevibacterium	lactofermentum	21800		2			1	
Brevibacterium	lactofermentum	21801						
Brevibacicium	lactofernientum		B11470	20				
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Brevibacterium	lactofernichtum	21086					1	
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Brevibacterium	lactofermentum	- 21086		_			+	T
Brevibacterum	lactofemicutum	31269					1	
Brevibacterium	linens	9174		-			+	
Brevibacicium	linens	19391						
Brevibacterium	linens	8377					1	
Brevibacterum	paraffinolyticum				09[[			
Brevibacterium	spec.				-	11.73		
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	acctoacidophilum	acetoacidophilum	acctoglutamicum	acetoglutamicum	acetoglutamicum	acetoglutamicum	acetoglutamicum	acctophilum	ammoniagenes	ammoniagenes	fujiokense	glutamicum	glutanucum.	glutamicum	glutamicum	glutamicum	glutamicum														
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	Corynchacterium	Corymebacterium	Corynebacterium	Corynebacterium	Corynebacterium	Cornebacterium	Corynebacterium	Corynebacterium	Coryncbacterium	Corynebacterium	Colynebacterium	Corynebacterium	Corynchacterium	Coryncbacterium	Corynebacterium	Corynebacterium	Corynebacterium	Corynebacterium	Corymebacterium	Corynebacterium											

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glutamicum	glutamicum	glutanicum	glutanicum	glutamicum	glutamicum	glutamicum	glutamicum	glutamicum	glutaniicum	glutamicum	glutamicum	glutamicum	glutar	glutar	glutar	glutar	glutar	e lutar	glutar	gluta	E luta	Blita	gluta	glula	gluta	Bluta	g luta	elufa	Elufa Bufa	Π	Π
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glutamicum	glutamicum	glutamicum	glutamicum	glutamicum	glutanicum	glutamicum	glutamicum	glutamicum	glutamicum	glutamicum	glutanicum	glutamicum	glutanicum	glutamicum	glutamicum	elutamicum	glutamicum	glutamicum	ilium	nitrilophilus	spcc.	spec.	spec.	spec.	spec.	spec.	spec	spec.	spec.	spcc.	spec.
	П			П	Γ	Π		T	П			Π	Г	1	Т	T	T	1	Т	1	Τ	Π		Т	Т	П		T	T	L	
Corynebacterium	Corynebacterium	Corynebacterium	Corynchacterium	Corynebacterium	Corynebacterium	Corynebacterium	Corynebacter ium	Cornebacterium	Corynebacterium	Corynebacterium	Corynebackerium	Corynebacterium	Coryncbacterium	Cormebacterium	Convietacterium	Corvachacterium	Corvnebacterium	Corynchacterium	Corynchacterium	Cornebacterum	Corynebacterium	Corynebacterium	Corynebacterium	Corynchacterium	Connebacterium	Correbacterium	Corynchacterium	Corynebacterium	Corynebacterium	Connebacterium	Corynebacterium
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ATCC: American Type Culture Collection, Rockville, MD, USA

FERM: Femientation Research Institute, Chiba, Japan

NRRL: ARS Culture Collection, Northern Regional Research Laboratory, Peoria, IL, USA

CECT: Coleccion Espanola de Cultivos Tipo, Valencia, Spain

NCIMB: National Collection of Industrial and Marine Bacteria Ltd., Aberdeen, UK

CBS: Centrastbureau voor Schimmetcultures, Baam, NL.

NCTC: National Collection of Type Cultures, London, UK

DSMZ: Deutsche Sammlung von Mikrooiganismen und Zellkulturen, Braunschweig, Germany

For reference see Sugawaia, H et al (1993) World directory of collections of cultures of microorganisms. Bacteria, fungi and yeasts (4th edn), World sederation

for culture collections would data center on microorganisms, Saimata, Japen.

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>>RXA01849-amino acid sequence (1-450, translated) 150 residues

LPGVELPDLI LSQIAWLCAR IEVDGMRADL VITRTALAHA AWAGRTVVTE EDVEIAARLA LPHRRRNPF DAPEMEERKL QETLQEARDF FKDNEDKGPA AKITDEETGA EAFTDTDNPT EEDGLQGTAQ AKAQTTGKVG TAGSGDPFRS

>RXA01849-nucleotide sequence B: coding region

CTGCCTGGTGTGGAGCTGCCGGATCTGATCTTGTCGCAGATTGCGTGGTTGTGTGCACGTATTGAAGTCGACGGTAT GCGCGCTGACCTGGTGATCACGCGTACCGCACTTGCTCACGCCGCGTGGGCTGGACGCACTGTGGTTACGGAAGAAG ACGTGGAGATCGCAGCTCGCCTAGCGTTGCCGCACCGCCGTCGCCGTAATCCTTTCGATGCTCCAGAAATGGAGGAG CGCAAGCTTCAGGAAACCCTGCAGGAAGCTCGGGACTTCTTCAAAGACAATGAAGATAAAGGACCTGCCGCCAAGAT CACCGATGAGGAAACCGGTGCAGAGGCCTTTACCGATACCGACAATCCCACCGAGGAAGACGGTCTGCAAGGAACTG CGCAGGCGAAGGCGCAGACTACTGGAAAGGTACTGCCGGATCCGGCGACCCCTTTCGCTCC

>RXA01849-nucleotide sequence C: downstream TAGGCATTTGCGCCTGGCGTCCA

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990662 ····· O.Z., 0050/50169

>>RXA01848-amino acid sequence

(1-744, translated) 248 residues

MGEEDSTPGR RSKAYSRQGA DVRPMKGGHG INLVGTLMAA TERGANIVEG VVDFRPTDLR GSLRRGREAN LIVFVVDTSG SMAARSRVRA VTGTITSMLN DAYQRRDKVA VIAVNGNKPT LVLNPTNSVE QAQQKLKDMP MGGRTPLAEG LLMAKDLMAR ELRKEPGRRA ILMVMTDGQD TSDAGEAGIA TAAETVVKSR LSGNVVIDCE GRLKVRKERA GVLAEMLGGV CVRLRDLNSE HIKMVINA

>RXA01848-nucleotide sequence A: upstream

>RXA01848-nucleotide sequence B: coding region

>RXA01848-nucleotide sequence C: downstream TAGACAACCAGAGTGAGGGTTTC

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.>>RXA01691-amino acid sequence

(1-567, translated) 189 residues

MASQQIRYPF SAVVGQDELR LALILTAISP RIGGVVIRGE KGTAKTTTVR AFAGLLGDAP LVNLPLGSTE DRVVGSLNME TVLTTGRAEY QPGLLAQADG GVLYVDEVNL LADHLVDALL DAAASGRVSI ERDGISHSSP ANFVLVGTMN PEEGELRPQL LDRFGLAVDV AASTNPEVRV EIIRRRLDF

>RXA01691-nucleotide sequence A: upstream

AAAACCTTAAGTTGGGTGGTTAAACCCACTAAGGTCTCACTTTATGGATGTGCCAGGTCACACAAAAAATCTCAAGAAAACTCACATTAAAGGACAGTA

>RXA01691-nucleotide sequence B: coding region

### Claims

- 1. An isolated nucleic acid molecule from Corynebacterium glutumicum encoding an HA protein, or a portion thereof.
- 2. The isolated nucleic acid molecule of claim 1, wherein said nucleic acid molecule encodes an HA protein involved in the production of a fine chemical.
- 3. An isolated Corynehacterium glutamicum nucleic acid molecule selected from the group consisting of those sequences set forth in Appendix A, or a portion thereof.
  - 4. An isolated nucleic acid molecule which encodes a polypeptide sequence selected from the group consisting of those sequences set forth in Appendix B.
- 15 5. An isolated nucleic acid molecule which encodes a naturally occurring allelic variant of a polypeptide selected from the group of amino acid sequences consisting of those sequences set forth in Appendix B.
- 6. An isolated nucleic acid molecule comprising a nucleotide sequence which is at least 50% homologous to a nucleotide sequence selected from the group consisting of those sequences set forth in Appendix A, or a portion thereof.
  - 7. An isolated nucleic acid molecule comprising a fragment of at least 15 nucleotides of a nucleic acid comprising a nucleotide sequence selected from the group consisting of those sequences set forth in Appendix A.
  - 8. An isolated nucleic acid molecule which hybridizes to the nucleic acid molecule of any one of claims 1-7 under stringent conditions.
- 30 9. An isolated nucleic acid molecule comprising the nucleic acid molecule of any one of claims 1-8 or a portion thereof and a nucleotide sequence encoding a heterologous polypeptide.
  - 10. A vector comprising the nucleic acid molecule of any one of claims 1-9.
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  11. The vector of claim 10, which is an expression vector.

- 12. A host cell transfected with the expression vector of claim 11.
- 40 13. The host cell of claim 12, wherein said cell is a microorganism.
  - 14. The host cell of claim 13, wherein said cell belongs to the genus Corynebucterium or Brevibucterium.
- 15. The host cell of claim 12, wherein the expression of said nucleic acid molecule results in the modulation in production of a fine chemical from said cell.

16. The host cell of claim 15, wherein said fine chemical is selected from the group consisting of: organic acids, proteinogenic and nonproteinogenic amino acids, purine and pyrimidine bases, nucleosides, nucleotides, lipids, saturated and unsaturated fatty acids, diols, carbohydrates, aromatic compounds, vitamins, cofactors, and enzymes.

- 17. A method of producing a polypeptide comprising culturing the host cell of claim 12 in an appropriate culture medium to, thereby, produce the polypeptide.
- 10 18. An isolated HA polypeptide from Corynebacterium glutamicum, or a portion thereof.

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- 19. The polypeptide of claim 18, wherein said polypeptide is involved in the production of a fine chemical production.
- 20. An isolated polypeptide comprising an amino acid sequence selected from the group consisting of those sequences set forth in Appendix B.
- 21. An isolated polypeptide comprising a naturally occurring allelic variant of a
   polypeptide comprising an amino acid sequence selected from the group consisting of those sequences set forth in Appendix B, or a portion thereof.
  - 22. The isolated polypeptide of any of claims 18-21, further comprising heterologous amino acid sequences.
- 23. An isolated polypeptide which is encoded by a nucleic acid molecule comprising a nucleotide sequence which is at least 50% homologous to a nucleic acid selected from the group consisting of those sequences set forth in Appendix A
- 30 24. An isolated polypeptide comprising an amino acid sequence which is at least 50% homologous to an amino acid sequence selected from the group consisting of those sequences set forth in Appendix B.
- 25. A method for producing a fine chemical, comprising culturing a cell containing a vector of claim 12 such that the fine chemical is produced.
  - 26. The method of claim 25, wherein said method further comprises the step of recovering the fine chemical from said culture.
- 27. The method of claim 25, wherein said method further comprises the step of transfecting said cell with the vector of claim 11 to result in a cell containing said vector.
- 28. The method of claim 25, wherein said cell belongs to the genus Corynebacterium or Brevibacterium.
  - 29. The method of claim 25, wherein said cell is selected from the group consisting of: Corynebacterium glutamicum, Corynebacterium herculis, Corynebacterium lilium, Corynebacterium acetoacidophilum, Corynebacterium acetoglutamicum,

Corynebucierium ucetophilum, Corynebacterium ammoniagenes. Corynebacterium fujiokense, Corynebacterium nitrilophilus. Brevibacterium ammoniagenes, Brevibacterium butanicum, Brevibacterium divaricatum. Brevibacterium flavum, Brevibacterium healii, Brevibacterium ketoglutamicum, Brevibacterium ketosoreductum, Brevibacterium lactofermentum, Brevibacterium linens, Brevibacterium paraffinolyticum, and those strains set forth in Table 3

- 30. The method of claim 25, wherein expression of the nucleic acid molecule from said vector results in modulation of production of said fine chemical.
- 31. The method of claim 25, wherein said fine chemical is selected from the group consisting of: organic acids, proteinogenic and nonproteinogenic amino acids, purine and pyrimidine bases, nucleosides, nucleotides, lipids, saturated and unsaturated fatty acids, diols, carbohydrates, aromatic compounds, vitamins, cofactors, and enzymes.
  - 32. The method of claim 25, wherein said fine chemical is an amino acid.
- 33. The method of claim 32, wherein said amino acid is drawn from the group consisting of: lysine, glutamate, glutamine, alanine, aspartate, glycine, serine, threonine, methionine, cysteine, valine, leucine, isoleucine, arginine, proline, histidine, tyrosine, phenylalanine, and tryptophan.
- 34. A method for producing a fine chemical, comprising culturing a cell whose genomic 25 DNA has been altered by the inclusion of a nucleic acid molecule of any one of claims 1-9.

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